Radiation-Induced Brain Structural and Functional Abnormalities in Presymptomatic Phase and Outcome Prediction

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Abstract: Radiation therapy, a major method of treatment for brain cancer, may cause severe brain injuries after many years. We used a rare and unique cohort of nasopharyngeal carcinoma patients with normal-appearing brains to study possible early irradiation injury in its presymptomatic phase before severe, irreversible necrosis happens. The aim is to detect any structural or functional imaging biomarker that is sensitive to early irradiation injury, and to understand the recovery and progression of irradiation injury that can shed light on outcome prediction for early clinical intervention. We found an acute increase in local brain activity that is followed by extensive reductions in such activity in the temporal lobe and significant loss of functional connectivity in a distributed, large-scale, high-level cognitive function-related brain network. Intriguingly, these radiosensitive functional alterations were found to be fully or partially recoverable. In contrast, progressive late disruptions to the integrity of the related far-end white matter structure began to be significant after one year. Importantly, early increased local brain functional activity was predictive of severe later outcome.

Zhongxiang Ding, Han Zhang, and Xiao-Fei Lv contributed equally to this work.

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tumor patients limits the follow-up time for the investigation of the post-RT brain injury at the time when no irradiation-induced brain abnormalities have already been detected [Wong and Van der Kogel, 2004]. An early, presymptomatic detection of the post-RT brain injury at the time when no magnetic resonance (MR)-visible lesions have developed is of great importance for early prevention and mitigation of these complications [Gondi et al., 2010].

Investigation of the brain injury in different stages after completion of RT is quite interesting to the neuroscience community, because of the complex time-evolving interplay between brain recovery, plasticity, and degeneration. However, previous studies on brain tumor patients have two major constraints. First, short survival time of brain tumor patients limits the follow-up time for the investigation of late irradiation-induced brain abnormalities [Chapman et al., 2012; Nagesh et al., 2008]. Second, the residual brain lesions or recurrences that may not be easily differentiated from the RT-induced injuries [Chan et al., 1999]. In contrast, nasopharyngeal carcinoma patients constitute an ideal cohort for such a study. Nasopharyngeal carcinoma is a malignant tumor originating in the nasopharynx [Chang and Adami, 2006]; RT is a routine clinical treatment [Brennan, 2006] and may inevitably result in irradiation of the normal brain tissue [Chen et al., 2011]. More than 80% of the RT patients survive for five years or longer [Zhou et al., 2013]; long enough for the study of the late effects of RT. However, almost all studies using nasopharyngeal carcinoma patients have focused on MR-visible brain injury at the specific brain regions (e.g., temporal lobe necrosis), which usually appears many years after RT [Chan et al., 2003; Chen et al., 2011; Chong et al., 2000; Su et al., 2012, 2013; Tang et al., 2014; Tsai et al., 2014; Ye et al., 2012; Zeng et al., 2015; Zheng et al., 2015; Zhou et al., 2013, 2014] at the stage when brain damage has already been irreversible. Therefore, the recovery ability is quite limited, and the reverse of the impaired cognitive function is usually impossible [Cheung et al., 2003].

We recently began to investigate RT-induced brain injury in nasopharyngeal carcinoma patients at its presymptomatic stage, at which time the patients still have normal-appearing MR or CT images [Wang et al., 2012; Xiong et al., 2013]. These studies, together with only a few other presymptomatic studies on brain tumor patients [Chapman et al., 2012; Matulewicz et al., 2006; Nagesh et al., 2008], focused only on the structural or morphometric changes in the brain white matter. However, these results have largely been inconsistent and have lacked spatial specificity due to the region of interest (ROI)-based analysis strategy. In addition, for these brain tumor studies, inhomogeneous cohorts with different tumor location, pathology and size resulted in different irradiated areas, while the craniotomy and tumor recurrence could have introduced additional confounding effect. However, the most significant issue for these studies is that different predefined ROI placements introduced tremendous variation in the measuring results and led to such inconsistent conclusions. For example, Xiong et al. [2013] calculated diffusion tensor imaging (DTI) metrics in a few preset ROIs within the temporal lobe. They found that the temporal lobe white matter was affected immediately after RT but recovered one year later. However, in another study, fractional anisotropy (FA) values were found to remain abnormally low after one year [Wang et al., 2012]. In a more recent study, Chen et al. [2015] found an acute decrease in FA within 3 months after RT in the anterior temporal lobe, but no data were available for the study of late responses. The above findings, however, were not consistent with those from the brain tumor studies, in which different white matter areas, such as the parahippocampal cingulum bundles [Chapman et al., 2012] and the genu and splenium of the corpus callosum [Nagesh et al., 2008] were assessed and revealed a gradual course towards abnormality, which again indicates progression of structural abnormalities. In addition to DTI, both our group [Wang et al., 2012; Xiong et al., 2013] and another group [Chen et al., 2014] have used magnetic resonance spectroscopy

**INTRODUCTION**

Radiation therapy (RT) is a longstanding and routine clinical treatment for cancer. During cranial RT, normal brain tissue along the RT pathway and in proximity to tumors is inevitably irradiated, causing not only transient and reversible abnormalities but also progressive and irreversible late toxicities such as brain tissue necrosis [Acker et al., 1998; Bowen et al., 1996; Chan et al., 2009b; Haddy et al., 2011; Lo et al., 1992; New, 2001]. RT-induced mild cognitive impairment and/or neuropsychiatric disorders also significantly decrease the quality of life [Greene-Schloesser and Robbins, 2012; Khong et al., 2006; Tang et al., 2012; Warrington et al., 2013]. Despite carefully designed irradiation delivery, RT may still result in such sequelae; but our understanding of the large-scale neuromechanism of the post-RT brain injury is still insufficient [Wang et al., 2010; Wong and Van der Kogel, 2004]. An early, presymptomatic detection of the post-RT brain injury at the time when no magnetic resonance (MR)-visible lesions have developed is of great importance for early prevention and mitigation of these complications [Gondi et al., 2010].

Investigation of the brain injury in different stages after completion of RT is quite interesting to the neuroscience community, because of the complex time-evolving interplay between brain recovery, plasticity, and degeneration. However, previous studies on brain tumor patients have two major constraints. First, short survival time of brain tumor patients limits the follow-up time for the investigation of late irradiation-induced brain abnormalities [Chapman et al., 2012; Nagesh et al., 2008]. Second, the residual brain lesions or recurrences that may not be easily differentiated from the RT-induced injuries [Chan et al., 1999]. In contrast, nasopharyngeal carcinoma patients constitute an ideal cohort for such a study. Nasopharyngeal carcinoma is a malignant tumor originating in the nasopharynx [Chang and Adami, 2006]; RT is a routine clinical treatment [Brennan, 2006] and may inevitably result in irradiation of the normal brain tissue [Chen et al., 2011]. More than 80% of the RT patients survive for five years or longer [Zhou et al., 2013]; long enough for the study of the late effects of RT. However, almost all studies using nasopharyngeal carcinoma patients have focused on MR-visible brain injury at the specific brain regions (e.g., temporal lobe necrosis), which usually appears many years after RT [Chan et al., 2003; Chen et al., 2011; Chong et al., 2000; Su et al., 2012, 2013; Tang et al., 2014; Tsai et al., 2014; Ye et al., 2012; Zeng et al., 2015; Zheng et al., 2015; Zhou et al., 2013, 2014] at the stage when brain damage has already been irreversible. Therefore, the recovery ability is quite limited, and the reverse of the impaired cognitive function is usually impossible [Cheung et al., 2003].

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(MRS) to measure brain tissue metabolism and neuronal function in post-RT, normal-appearing nasopharyngeal carcinoma patients, but this modality is hampered by heavy noise and poor spatial resolution [Jansen et al., 2006], as well as the same ROI-definition problem. More advanced brain functional study using functional magnetic resonance imaging (fMRI) has been largely neglected. Collectively, due to the use of these oversimplified, less detailed, and potentially biased methods of analysis, precise localization of RT-induced brain structural and functional changes in the presymptomatic stage remain largely unknown. A whole-brain, voxel-wise exploration is highly desirable [Lv et al., 2014] to provide a better understanding of how the brain evolves in the natural post-RT course, during which recovery and progression can dynamically interact.

The aim of this study is to explore RT-induced brain functional abnormality in the presymptomatic stage in a voxel-wise manner by taking advantage of relatively high spatial and temporal resolution of fMRI. An additional aim is to try to understand the relationship between functional and structural abnormalities in a long post-RT period, with which we can better characterize the structurally and functionally evolutionary processes of the human brain after the completion of RT. It is more clinically important to investigate presymptomatic stage than symptomatic or clinical stage because early alterations, especially those that can be detected by more sensitive imaging and analysis techniques, can potentially be used in the future to predict the outcomes many years later, thereby allowing adequate time for neuroprotective or other preventive treatment. In this article, we report our initial finding based on a prospective, follow-up, pseudo-longitudinal cohort study. To this end, we divided different normal-appearing nasopharyngeal carcinoma patients into four groups: a baseline group (prior to RT, G1) and three post-RT groups (G2, G3, and G4). The patients in the latter three groups were scanned during different post-RT periods ranging from the acute to the delayed response stage (0-6, 6-12, and >12 months for G2, G3, and G4, respectively). For each group, multimodal brain imaging, including rs-fMRI and DTI were conducted for probing both brain function and structure. These patients were similarly treated with RT, regularly followed up with clinical MRI, until some of them showed MRI-visible RT-induced injuries (e.g., temporal lobe necrosis), based on which our detected early imaging biomarker were validated.

Specifically, the brain function was measured by resting-state fMRI (rs-fMRI) using both localized brain activity and inter-regional co-activity (functional connectivity, or FC) as a natural resting state is helpful for detecting aberrance in inherent brain spontaneous activities and their inter-regional coactivity. Based on previous RT-induced focal lesion studies, we believed that, the brain spontaneous activity at the region nearby or in the path of the irradiation (i.e., the near-end brain regions) could show early abnormalities. Previous clinical studies have shown that fractional amplitude of low-frequency fluctuation (fALFF) [Zang et al., 2007; Zou et al., 2008] is a reliable and sensitive metric for such purpose. Therefore, we used fALFF to detect RT-induced local brain functional abnormalities. We also investigated the possible reason of the RT-induced mild cognitive impairment, a common side effect of RT [Greene-Schloesser and Robbins, 2012], using FC based on temporally synchronized brain spontaneous activities because the previous studies have shown that FC or resting-state brain functional network is one of the important neuromechanisms for retaining healthy and normal cognitive functions and responsible for abnormal cognitive abilities or mental disorders [Buckner, 2013]. According to the literature, it is the default mode network (DMN) that has often been found to be mostly related to the mild cognitive impairment due to Alzheimer’s disease (AD), Parkinson’s disease (PD) and other diseases (e.g., Type-II diabetes), we thus also focused on the FC in the DMN and hypothesized that both near- and far-end regions in the DMN could be affected by RT. Moreover, the DMN includes hippocampus and other temporal regions that have been extensively reported in previous post-RT brain injury studies; therefore, DMN is an ideal functional network for us to investigate. Another potential RT-influence on the far-end regions could be the white matter structures that link to the affected near-end temporal regions. We thus use FA based on DTI to investigate the microstructural integrity of the white matter structure to further support our functional imaging-based findings.

Collectively, the hypotheses were as follows: (1) RT affects both localized functioning and functional integration in the near-end temporal lobe, particularly the hippocampus, at the acute and early response stages; (2) FC to the far-end cognitive function-related regions such as posterior cingulate cortex (PCC) in the DMN will also be affected later as some of the DMN connectivities are along the irradiation pathway; and (3) regardless of whether the brain function changes in grey matter could recover (or be plastic) or not, long-term degenerative injuries in white matter could manifest as progressively developing FA abnormalities. The innovation of our study is three-fold. First, this is the first multimodal voxel-based brain structural and functional study on the post-RT brain abnormalities using MR-normal-appearing patients with non-brain tumors. Second, this is also the first study that investigates the potential relationship between brain functional and structural changes after RT. Third, this is the first study using rs-fMRI to detect early sign of RT injury which is found to be dose dependent. Based on our results, we, for the first time, tentatively plot a recovery/progression course with respect to the post-RT brain functional and structural changes, which highlight the ability of human brain under continuous focal attacks. We also propose a preventive model that could serve as a guideline for future clinical practices to prevent or reduce the probability of RT-induced severe brain injury.
TABLE I. Demographic and clinical information of the subjects

<table>
<thead>
<tr>
<th>G1</th>
<th>G2</th>
<th>G3</th>
<th>G4</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td># Subjects (M/F)</td>
<td>25 (21/4)</td>
<td>24 (20/4)</td>
<td>19 (12/7)</td>
<td>19 (11/8)</td>
</tr>
<tr>
<td>Age (mean ± SD)</td>
<td>45.08 ± 8.56</td>
<td>44.83 ± 9.07</td>
<td>41.53 ± 9.63</td>
<td>45.00 ± 12.57</td>
</tr>
<tr>
<td>T-classification</td>
<td>2/4/14/5</td>
<td>1/6/13/4</td>
<td>1/3/10/5</td>
<td>1/6/11/1</td>
</tr>
<tr>
<td>N-classification</td>
<td>3/14/6/2</td>
<td>2/11/8/3</td>
<td>2/9/6/2</td>
<td>3/7/6/3</td>
</tr>
<tr>
<td>M-classification</td>
<td>25/0/0/0</td>
<td>24/0/0/0</td>
<td>19/0/0/0</td>
<td>19/0/0/0</td>
</tr>
<tr>
<td>Tumour Overall Stage</td>
<td>1/5/13/6</td>
<td>1/5/11/7</td>
<td>1/3/8/7</td>
<td>1/6/7/5</td>
</tr>
<tr>
<td>Scan Time (mo) Pre-RT</td>
<td>0–6 mo</td>
<td>6–12 mo</td>
<td>&gt; 12 mo</td>
<td>—</td>
</tr>
<tr>
<td>RT-injury Stage</td>
<td>Baseline</td>
<td>AC, ED</td>
<td>LD</td>
<td>LD</td>
</tr>
<tr>
<td>Spec Scan Time (mo)</td>
<td>3.08±1.53</td>
<td>10.89±1.56</td>
<td>33.37±19.26</td>
<td>0.05f</td>
</tr>
<tr>
<td>#Subjects 2D-CRT/IMRT</td>
<td>—</td>
<td>16/8</td>
<td>10/9</td>
<td>17/2</td>
</tr>
<tr>
<td>#Subjects Chemo</td>
<td>—</td>
<td>23</td>
<td>18</td>
<td>18</td>
</tr>
</tbody>
</table>

*M: male; F: female; mo: months; scan time: the mean ± SD of specific time interval in months between completion of radiotherapy and MR scan; pre-RT: prior to radiotherapy; spec scan time: the mean ± SD of specific time interval in months between completion of radiotherapy and MR scan; IMRT: intensity-modulated radiation therapy; 2D-CRT: two-dimensional conventional radiation therapy; Chemo: chemotherapy. AC: acute stage; ED: early delayed stage; LD: late-delayed stage; G1–4: groups 1–4. T-classification describes the size of the primary tumor and whether it has invaded nearby tissue and has four levels (we describe the number of patients who are, from mild to severe, T1/T2/T3/T4). N-classification describes nearby lymph nodes that are involved (four levels, N0/N1/N2/N3, from mild to severe). M-classification defines distant metastasis (no or yes, M0/M1). Tumor overall stage (I, II, III, and IV) was obtained based on the T-, N-, and M-classification results. The number of patients at each of the four stages (I/II/III/IV) is listed.

MATERIALS AND METHODS

Subjects

Newly diagnosed, treatment-naïve nasopharyngeal carcinoma patients with no brain invasion were included. Tumor stage was assessed according to the guidelines reported previously [Li et al., 2014]. The subject inclusion criteria for this study were: (1) newly diagnosed with non-keratinizing, undifferentiated nasopharyngeal carcinoma by histopathology; (2) hospitalized between 2005 and 2012; (3) no intracranial invasion; (4) clinically silent lesions; (5) neither a brain tumor nor distant metastases; (6) between 18 and 70 years of age; (7) no drinking or smoking habits; (8) no neurological or psychiatric diseases or other major medical issues; and (9) being right handed. Clinically routine MRI, including thick-slice T2- and T1-weighted images and T2 fluid-attenuated inversion recovery images, were obtained to make sure that there was no brain invasion before or any visible lesions after RT. This initial visual screening was performed by two experienced diagnostic radiologists (XL and FX, with 9 and 5 years of experience, respectively) at the initial inclusion and was verified by another diagnostic radiologist (ZD, with 24 years of experience) before data processing. Two diagnostic radiologists (XL and FX) used the internationally recommended 7th edition of the Union for International Cancer Control-American Joint Committee on Cancer (UICC-AJCC)’s Tumor, Node, Metastasis (TNM) staging system for nasopharyngeal carcinoma stage classification. The patients’ TNM stage ranged from T1N0M0 to T4N2M0. All subjects underwent a detailed pretreatment evaluation, including a physical examination, nasopharyngeal fiberoptic endoscopy, MRI of the nasopharynx and neck, chest radiography, abdominal sonography, and a whole body bone scan, to exclude any associated diseases. A total of 90 patients (44.27 ± 9.79 years; range: 19–68 years; 23 females and 67 males) from Sun Yat-sen University Cancer Center were included. Based on the subsequent data processing results, three subjects were excluded due to excessive head motion. Finally, 87 subjects were included (44.22 ± 9.85 years; range: 19–68 years; 23 females). There were more males than females due to the significantly higher incidence of nasopharyngeal carcinoma in males [Chang and Adami, 2006]. There were no significant differences in age, gender, tumor TNM stage, or overall stage among the groups. For details, please see Table I. This study was approved by the Ethics Committee of Sun Yat-sen University Cancer Center. Written informed consent was obtained from the participants.

Treatments

Sixty-two of the patients in this study had been treated with RT as described earlier [Lv et al., 2014; Sun et al.,
We implemented a cross-sectional comparison strategy because the subjects were scanned only once. Based on the time between RT completion and MRI acquisition, the subjects were divided into four groups (G1, G2, G3, and G4; N = 25, 24, 19, and 19, respectively; see Table I). G1 comprised treatment-naive nasopharyngeal carcinoma patients (i.e., pre-RT baseline), whereas G2, G3, and G4 comprised patients who had completed RT 0–6 months (including acute and early delayed stages), 6–12 months (late-delayed stage), and >12 months (late-delayed stage) before MRI acquisitions, respectively, similar to our previous study [Wang et al., 2012].

### Data Acquisition

We used two noninvasive imaging modalities, rs-fMRI, and DTI. Rs-fMRI measures the blood oxygen level-dependent (BOLD) signal, which reflects spontaneous brain activity during the resting state. Rs-fMRI has the same good temporal and spatial resolution as traditional task-based fMRI, but it allows us to measure both regional brain activity and inter-regional co-activity (FC). DTI maps the diffusion of water molecules in the brain, thereby measuring the microstructural integrity of neuronal fibers in white matter. In addition, DTI-based diffusion tractography can trace major fiber bundles, the spatial substrate of the FC [Hagmann et al., 2006].

All the images were collected using a 3.0-T MR scanner (Siemens Magnetom Tim Trio, Erlangen, Germany). The imaging parameters were (1) rs-fMRI: echo-planar imaging sequence, acquisition matrix = 64 × 64, field of view = 230 × 230 mm², 40 axial slices, slice thickness = 3.59 mm, interslice gap = 0.73 mm, voxel size = 3.59 × 3.59 × 3.59 mm³, repetition time = 2,400 ms, echo time = 30 ms, and 240 time points (9 min 36 s); (2) high-resolution, three-dimensional T1-weighted structural MRI: magnetization prepared rapid acquisition gradient echo sequence, acquisition matrix = 256 × 256, field of view = 256 × 256 mm², 176 sagittal slices, no interslice gap, voxel size = 1 × 1 × 1 mm³, flip angle = 9°, echo time = 2.52 ms, inversion time = 900 ms, and repetition time = 1,900 ms; and (3) DTI:

### TABLE II. Dose (in mGy) of radiation therapy for the G2 and G3 subjects

<table>
<thead>
<tr>
<th>Group #</th>
<th>N</th>
<th>Min</th>
<th>Max</th>
<th>Mean</th>
<th>Min</th>
<th>Max</th>
<th>Mean</th>
</tr>
</thead>
<tbody>
<tr>
<td>G2</td>
<td>8</td>
<td>120.8±79.4</td>
<td>7,150.5±417.9</td>
<td>1,691.7±469.8</td>
<td>108.6±66.7</td>
<td>6,906.0±587.6</td>
<td>1,758.1±711.4</td>
</tr>
<tr>
<td>G3</td>
<td>9</td>
<td>146.2±86.2</td>
<td>6,831.7±550.9</td>
<td>1,750.1±616.3</td>
<td>132.2±67.0</td>
<td>6,806.2±626.2</td>
<td>1,695.9±591.6</td>
</tr>
<tr>
<td>G4</td>
<td>2</td>
<td>208.6–301.3</td>
<td>6,882.0–6,993.4</td>
<td>2,036.1–2,368.5</td>
<td>210.3–178.5</td>
<td>6,940.0–6,818.6</td>
<td>2,086.3–1,894.2</td>
</tr>
</tbody>
</table>

*Dose refers to the dose–volume statistics applied to each side of the temporal lobe in patients who were treated with the IMRT protocol (N = 19, see # subjects with dose). For the dose of Groups 2 and 3, both the mean ± SD and the range are given, while for Group 4, only the range is given due to the small sample size.

**p:** the P value generated by two-sample t test on the doses between G2 and G3. N: sample size. Left/right side: dose calculated on the left/right side of the temporal lobe; Min: the minimum dose within the RT target area in mGy; Max: the maximum dose in an area of at least 2 mm² within the RT target in mGy; Mean: the averaged dose in mGy across all units within the RT target area. G2: the group of subjects scanned 0–6 months after RT; G3: the group of subjects scanned 6–12 months after RT; G4: the group of subjects scanned more than 12 months after RT; RT: radiation therapy.
Schematic flowchart of this study. With fMRI data (A), both the fALFF (measuring local activity) and the seed correlation-based FC (measuring inter-regional co-activity) were calculated and compared between G1 and the other groups (B). With DTI data, a voxel-wise comparison of the FA (measuring white matter microstructural integrity) was conducted between G1 and the other groups (D). Fiber tracking (E) was performed to link local functional changes (G) and inter-regional functional integration changes (C) with white matter changes (F, H). [Color figure can be viewed at wileyonlinelibrary.com]

Figure 1.

**Local Activity: fALFF Analysis**

The rs-fMRI and high-resolution T1 data were preprocessed with DPARSF2.2 [Yan and Zang, 2010], a toolbox based on REST1.8 [Song et al., 2011] and SPM8 (http://www.fil.ion.ucl.ac.uk/spm) in Matlab 2013a (MathWorks, Inc., Natick, MA). For rs-fMRI, the first five (12 s) images were discarded to allow for signal stabilization. The remaining data were corrected for both slice timing and head motion. Subjects with head motion >3 mm or 3° were excluded from further analyses. The T1 image was first co-registered to the rs-fMRI data and then segmented using Diffeomorphic Anatomical Registration Through Exponentiated Lie Algebra. The deformation field was then applied to the fMRI data to register them to the standard MNI space. For rs-fMRI, the last five (12 s) images were discarded to allow for signal stabilization. The remaining data were corrected for both slice timing and head motion. Subjects with head motion >3 mm or 3° were excluded from further analyses. The T1 image was first co-registered to the rs-fMRI data and then segmented using Diffeomorphic Anatomical Registration Through Exponentiated Lie Algebra. The deformation field was then applied to the fMRI data to register them to the standard MNI space. The fMRI data were further resampled to a resolution of $3 \times 3 \times 3$ mm$^3$ and spatially smoothed using a 6-mm, full-width, half-maximum, isotropic Gaussian kernel. Based on the preprocessed rs-fMRI data, fALFF was calculated using the ratio of the summed BOLD signal to the total signal within a region of interest.

**Data Analysis**

We used both fALFF and FC in DMN from rs-fMRI to quantify RT-induced functional abnormalities at the near- and far-end regions. The temporal precedence of RT-induced abnormalities in brain function and structure was investigated, along with different post-RT phases, to assess the course of post-RT recovery and/or progression. A schematic flowchart is provided in Figure 1.
fluctuation power within the low-frequency (0.01–0.08 Hz) part of the spectrum to that within the entire frequency range [Zou et al., 2008]. Because fALFF is considered less sensitive to physiological noise and artifacts in the bottom brain, we used fALFF rather than the original amplitude of low-frequency fluctuation [Zang et al., 2007]. We first computed fALFF for every voxel in each subject, resulting in individual-level fALFF maps. These were then standardized into z-score maps by subtracting the mean voxel-wise fALFF obtained for the entire brain and dividing by the standard deviation. This standardization can improve subsequent statistical analysis and test–retest reliability [Zuo et al., 2010].

Seed-Based FC Analysis

In addition to the local activity, RT can affect FC and cause cognitive impairment. This is the first report of the use of FC in a study of post-RT brain injury. Seed-based correction, a reliable and widely adopted method in the rs-fMRI community [Shehzad et al., 2009], was used to calculate FC, which measures the synchronization of the BOLD signals from different voxels by calculating Pearson’s correlation coefficients. We chose seed-based FC rather than other more complex metrics for easy interpretation [Friston, 2011]. The preprocessed rs-fMRI data were further detrended to remove linear trends due to drift, and they were band-pass filtered (0.01–0.08 Hz) to remove extremely low- and high-frequency artifacts. In addition, the averaged BOLD signal within the masks of cerebrospinal fluid and white matter and the estimated head motion parameters (Friston 24-parameter model) were regressed out to remove additional artifacts [Yan et al., 2013]. Framewise displacement (FD), the micro-head motion that has been found to influence the FC estimation, was calculated based on the method in Power et al. [2012]; the data volumes with FD > 0.5 were discarded, together with one data volume before and two volumes after them. After the data censoring, we further checked the remained data volumes for each subject. All the remained 87 subjects had adequate (8.96 ± 0.78 min, longer than 4.5 min) data for reliable FC calculation [Wu et al., 2015]. There was no group difference in the number of data volumes remained among the four groups (one-way ANOVA, F-test, df = 3, 83, P > 0.1). In addition, we also calculated overall translation in the three directions using mean root-mean-square (RMS) [Van Dijk et al., 2012] and overall micro-head motion using mean FD [Power et al., 2012]; there was no group difference in either metrics (one-way ANOVA, F-test, df = 3, 83, P > 0.1). To investigate FC in the DMN, a seed in the PCC [Shehzad et al., 2009] was chosen for the calculation of FC. Specifically, a 6-mm sphere centered on the PCC (MNI coordinates: −6, −58, 28) was chosen as the seed region for the FC analysis because it produces a reliable DMN. Individual FC maps were generated by calculating Pearson’s correlation coefficients between BOLD signals at every brain voxel and those in the seed region. These FC maps were then transformed into z-score maps for statistical analysis by subtracting the averaged FC in the brain and dividing by the standard deviation of FC across all brain voxels.

White-Matter Microstructural Integrity: FA Analysis

FA is a scalar that describes the degree of anisotropy of a diffusion process; it has been widely used to measure the microstructural integrity (e.g., myelination) of white matter [O’Donnell and Pastorinak, 2015]. A reduction in the FA is usually regarded as indicating deterioration of white-matter structural connections [Roosendaal et al., 2010; Voets et al., 2012]. Previous ROI-based FA analysis may constrain the result in specific brain regions; thus, for the first time, we calculated voxel-wise FA for the entire brain. The DTI data were processed with PANDA 1.30 [Cui et al., 2013], a toolbox based on FSL (http://fsl.fmrib.ox.ac.uk/fsl), Diffusion Toolkit (http://trackvis.org/dtk), and TrackVis (http://www.trackvis.org). The details of DTI data processing are summarized as follows: (1) estimating the brain mask using the $b = 0$ s/mm$^2$ image and the bet command of FSL to remove nonbrain regions; (2) correction for the eddy current effect by registering multiple diffusion-weighted images to the $b = 0$ s/mm$^2$ image with affine transformation; and (3) calculating voxel-wise diffusion tensors and FA. Note that the individual FA maps were originally calculated in native space. To allow for comparison across subjects, the FSL non-linear image registration tool, fiirt, was used to register each subject’s FA image of native space to the FA template in the standard MNI space. The FA images of the standard space were then resampled to $2 \times 2 \times 2$ mm$^3$ and spatially smoothed using a 6-mm full-width half-maximum isotropic Gaussian kernel for statistical analysis.

Statistical Analysis

Statistical analyses were conducted with REST 1.8 for the rs-fMRI data and with SPM8 for the DTI data. The post-RT groups (G2, G3, and G4) were separately compared with the pre-RT group (G1) using separate two-tailed, two-sample t tests in the framework of a general linear model unless specifically indicated otherwise. Two covariates—age and gender—were added to the model during group comparisons to regress out potential nuisance effects. Different inclusive masks were used during the comparisons of different metrics according to both their biological meanings and our hypotheses. Comparisons of fALFF and FA were conducted within a gray matter mask and a white matter mask, respectively; FC was compared within a DMN mask. Specifically, for the fALFF, a whole-brain gray matter mask (applying a threshold of $>0.2$ to SPM8’s a priori gray-matter template) was used because gray-matter voxels are generally considered to have meaningful activity. For the FC, a DMN mask was generated by conducting a one-sample t test on the
individual z-maps of the FC using all 87 subjects. A binary map was generated by applying a threshold of $P < 1 \times 10^{-12}$, which was false-discovery rate corrected ($t_{66} > 8.6$). To test if the subject’s framewise head motion contributed to the group difference detected, we also included mean FD as a quantified head motion index as an additional covariate (in addition to age and gender) during the two-sample t tests, the result (not shown) was highly similar to that without head motion covariate. Therefore, framewise head motion was unlikely to affect our result of FC differences. For the voxel-wise comparison of the FA, a mask was generated by averaging the unsmoothed FA maps across all 87 subjects; the resulting FA map was binarized with a threshold of $>0.4$. For the FA comparisons, multiple comparison corrections were applied based on a Gaussian Random Field method in SPM8, a toolbox commonly used for voxel-based analysis of FA, with a single-tailed, voxel-level intensity threshold of $P < 0.05$ ($z > 1.65$) and an extension threshold of $P < 0.05$ (cluster size $>990$ voxels). Single-tailed tests were used because of the hypothesis of reduced FA after RT. For the fALFF/FC results, a commonly used threshold for rs-fMRI studies, $P < 0.05$ after AlphaSim correction as implemented in REST was applied ($i.e.$, two-tailed voxel-wise $P < 0.05$, randomization times $=10,000$, resulting in cluster sizes $>172$ voxels for fALFF and $>101$ voxels for FC due to the different masks used in the $t$ tests). The course of recovery/progression after RT was investigated by comparing the statistically significant changes in fALFF, FC, and FA across the three post-RT periods.

**DTI Tractography**

Regarding the relationship between post-RT brain functional and structural changes, we further tested whether the changes in local brain activity are related to the disrupted FC and the reduction of white matter microstructural integrity. To this end, the clusters with significant FA reduction from the analyses of DTI data were selected as ROIs. These ROIs were transformed from the standard to the native space based on the invert warping function in FSL, in which the non-linear transformation from a native FA image to the FA template was first inverted and then applied to the ROIs. Diffusion tractography was conducted to reveal the main white-matter fiber bundles that cross these ROIs using deterministic fiber tracking, as implemented by the *fimplicit* function in FSL. Specifically, whole-brain fiber tracking was conducted first, with an automatic FA mask and an angle threshold of 40°. During fiber tracking, four seeds were randomly defined at each voxel. The main fiber bundles that intersect with each ROI were selected with TrackVis. Spatial closeness between these fibers and the gray-matter functional changes (these functional ROIs were selected based on the statistically significant results from the comparisons of fALFF and FC and warped back to the native space) was assessed for each subject (Fig. 1). According to the hypotheses, the fiber tracts that pass the affected white-matter areas would connect to the functionally affected areas (i.e., the functional ROIs).

**Follow-Ups and Retrospective Inspections**

This is a prospective study consisting of the aforementioned MRI scans at the subject inclusion, and regular follow-up scans for all the subjects using clinical MRI to detect MRI-visible RT-induced severe brain injury, that is, temporal lobe necrosis. Specifically, for the patients with post-RT rs-fMRI and DTI data (those in G2, G3, and G4), we conducted continuous follow-up observations every 3 months with clinical imaging sequences (T2-weighted and T2-FLAIR imaging). The patients with abnormal MR intensities in the unilateral or bilateral temporal lobe were regarded as exhibiting temporal lobe necrosis. Some patients inevitably dropped out during the follow-ups. Based on whether temporal lobe necrosis occurred during 5 years after RT completion, we retrospectively inspected the earlier brain functional/structural changes in the pre-symptomatic stage to search for predictive biomarkers or early signs of severe RT injury.

**RESULTS**

**RT-Induced Local Brain Activity Changes**

Compared with the baseline pre-RT group (G1), subjects in G2 (0–6 months after RT) had increased fALFF in the left inferior temporal areas (Fig. 2A) (“G2–G1” is used hereafter for comparison between G2 and G1). G2–G1 also showed an fALFF increase in the superior frontal area and an fALFF decrease in the supplementary motor area. G3–G1 showed a significant extensive decrease in the fALFF in the dispersed right temporal area and part of the supplementary motor area (Fig. 2B), but the superior frontal area still showed an fALFF increase. G4–G1 showed decreased fALFF in only a few areas in the bottom of the brain, including the bilateral hippocampus, the parahippocampus and the temporal pole, and the thalamus, brainstem, and cerebellar areas (Fig. 2C). The frontal area still had increased fALFF, which extended to the inferior frontal gyrus. In Figure 2D, the major brain regions with abnormal local activity are delineated for better illustration of the fALFF changes over time. The details of these differences are listed in Table III. We also inspected whole-brain fALFF differences with and without thresholds, and also found dynamic fALFF changes within and beyond the temporal lobe.

**RT-Induced FC Changes in DMN**

The seed-based correlation showed a complete DMN, consistent with previous studies; in particular, both the lateral and medial temporal cortices were included. These
lateral and medial temporal regions are the near-end regions exposed to the direct irradiation. The other DMN regions are thus far-end regions that could have abnormal connections to the near-end regions. There were no significant differences for G2–G1 and G4–G1. However, G3–G1 revealed a reduced FC with the PCC in various DMN regions (Fig. 3A), including the bilateral hippocampus, the parahippocampus, and other nontemporal lobe regions such as the medial prefrontal cortex, the right PCC/precuneus, and the left inferior parietal lobule (detailed in Table IV). Figure 3B shows the typical slices illustrating the FC changes within the DMN at different post-RT stages; several key regions of the DMN, including not only the hippocampus but also several frontal and parietal areas, were found to lose synchronization with the PCC. Whole-brain inspection without thresholds indicated that the above result still holds. Of note, we believed that head motion was not likely to contribute to the FC group differences because there was no group difference in either absolute or relative head motions, and because there was no visible differences in the FC different result when head motion were included as an additional covariate.

RT-Induced White-Matter Microstructural Integrity Changes

Within an FA mask in the putative white matter, there was a significant reduction in the FA value in G4–G1 (Fig. 4A) but not in G3–G1 or G2–G1. Specifically, after 12 months of the follow-ups from the completion of RT, the FA in the bilateral splenium of the corpus callosum was reduced compared to the pre-RT level (Table IV). Figure 4B clearly shows extensive FA reduction at >12 months, whereas at both 0–6 months and 6–12 months, the FA was quite stable. There was no significant post-RT late-delayed reduction in the FA within the temporal lobe white matter. The group differences in the entire white-matter mask without thresholds were also inspected and such a dynamic trend still held.

Recovery/Progression Course and “Preventive Model”

Based on the statistical analysis reported above, the course of schematic recovery/progression could be plotted (Fig. 5A). The involved regions included zones traditionally believed to be RT injury related (within the temporal lobe) and zones outside them. In the temporal lobe, the time-evolving pattern of local brain activity first increased, then decreased, and finally recovered. The most severe influence on local activity occurred in the late-delayed period (6–12 months), but there was an initial abnormal increase (<6 months). The FC with the PCC seed decreased at 6–12 months but recovered after 12 months. Functional recovery was accompanied by a gradual reduction in white-matter integrity in the corpus callosum after 12 months. Based on this temporal sequence, we derived a hypothesis for the prevention or delay of late effects,
which we call the preventive model. The hypothesis under this model is that, in future, by using early brain function monitoring with fALFF, one could implement protective treatment in time that may reduce the possibility of later FC reduction and white-matter injury.

**Predictive Biomarkers/Early Sign of RT Injury**

To validate our proposed preventive model, we further test if the early abnormality in local fALFF could predict severe late-delayed brain injury. Based on the follow-up results, five subjects in G2 (0–6 months after RT), one subject in G3 (6–12 months after RT), and two subjects in G4 (>12 months after RT) developed temporal lobe necrosis (regarded as “bad” outcome). Those who still have normal-appearing brain imaging result after 5 years of the follow-ups were categorized as “good” outcomes. We focused on the G2 group because this group has relatively larger sample size with the validated good/bad outcomes. In the G2 group, there were five subjects with >5 years’ follow-ups who still had no abnormal appearance in any part of the brain (as evaluated by clinical scans). As we have found an early abnormal increment of fALFF in this group of subjects compared with baseline pre-RT subjects in the G1 group, the averaged fALFF value within an ROI centered at the peak coordinates of G2–G1 fALFF group difference map in the temporal lobe was calculated for each subject in G1 and G2; these values were plotted in Figure 5B. As we expected, the five subjects with bad outcome had greater early fALFF increases in the left inferior temporal area than the five subjects with better outcomes. The median level of fALFF (normalized fALFF = −0.44) for the G2 subjects, as shown in Figure 5B, was able to successfully separate all the five subjects with bad outcomes from all the five subjects with good outcomes. Therefore, this line can be regarded as a “red line” for future RT treatment with continuous brain activity monitoring. The two subjects without later necrosis displayed fALFF values even lower than the median level of G1 subjects; these two subjects could be considered much safer.

**Relationship Between Functional and Structural Changes**

DTI tractography in a randomly selected subject showed a close relationship between the earlier reduction in temporal-lobe local brain activity and the later impairment of white matter integrity; this relationship was reflected by the fiber bundles connecting the two types of clusters (Fig. 6, upper panel). The fibers consist of a subset of callosal fibers that project to the temporal lobe (tapetum) and to the parietal and occipital lobes (forceps posterior); they may include the associated fibers projecting from the occipital lobe to the frontal and temporal lobes (parts of the inferior longitudinal fasciculus and the inferior fronto-
RT-induced DMN functional connectivity changes. Only G3 – G1 (6–12 months after RT vs pre-RT) revealed a significant FC reduction within the DMN ($t > 2.02$, $P < 0.05$, cluster size $> 101$, corrected by AlphaSim) (A). The unthresholded results for each post-RT stage on typical slices of the DMN are also shown (B), with a significant whole network FC reduction at 6–12 months after RT. The results were rendered on the averaged T1 images across all subjects. The brain slices are shown in radiological convention (left is right, right is left). Details of the results are summarized in Table IV. [Color figure can be viewed at wileyonlinelibrary.com]

Because patients in G3 had both decreased fALFF and decreased FC in the temporal lobe 6–12 months after RT, we performed a post hoc inspection and found that these results overlapped in the right hippocampus. There was a significant correlation between the fALFF and FC values in the right hippocampal ROI across all 87 subjects ($r = 0.23$, $P = 0.03$) (Fig. 7A). This means that if the fALFF in the hippocampus decreased, the FC between the PCC and the hippocampus also decreased.

### Dose Dependence

Although the local brain activity seems to occur earlier in the first 6 months and be able to predict later severe consequences, we regarded this increased fALFF as a local response to irradiation. As the inter-regional FC pathway is more related to neurodegenerative process in the far-end zone.
regions, and such a pathway could be more susceptible to RT injury due to its more exposure to irradiation; we proposed that the FC is more sensitive to irradiation dosage.

To test this hypothesis, we correlated the FC strength in the bilateral hippocampal ROIs (with FC reductions in G3–G1) with the RT doses to the temporal lobe across nine G3

Figure 4.

FA reductions 12 months or longer from the completion of RT. Only G4–G1 reached statistical significance (Gaussian Random Field-based correction, cluster-wise \( P < 0.05 \), voxel-wise \( P < 0.05 \), single-tailed statistical test as implemented in SPM8) (A). FA changes compared with G1 (pre-RT) at all three stages are shown without thresholds in several typical slices. The underlying image is the averaged T1 image across all subjects. The brain slices are shown in radiological convention (left is right, right is left). Details are summarized in Table IV. [Color figure can be viewed at wileyonlinelibrary.com]

Hypothesized model of different recovery/progression courses after RT with and without early intervention (A) and potential predictive biomarkers with early fALFF increments (B). In our hypothesized recovery/progressive model, a local temporal lobe brain activity increment preceded later default-mode FC and even later white matter integrity reductions. This model is plotted by the solid curves. We also proposed a preventive model, shown by the dotted curves; in this model, if the earlier local brain activity changes could be continuously monitored and carefully controlled by neuroprotective treatment, severe later white matter integrity loss and temporal lobe necrosis might be prevented. In (B), the early abnormal fALFF increment was plotted with the later outcomes (necrosis, indicating bad outcomes; or no necrosis after 5 years, indicating good outcomes) to evaluate the prognostic value of our proposed early biomarker and to further validate our preventive model. Specifically, based on the peak coordinates of G2–G1 group differences in fALFF in the left inferior temporal lobe (\(-45 -9 -39\), see Table III), a sphere ROI was generated (radius = 6 mm). Mean fALFF (in \( z \) score) across voxels in this ROI was extracted for each subject in G1 (pre-RT) and G2 (0–6 months after RT). Subjects showing clear temporal lobe necrosis during follow-up are shown in red; those whose brains still appeared normal are shown in blue. Subjects with unclear outcomes due to dropout are shown in gray. The median fALFF value for G2 subjects \((z = -0.44)\) delineates a warning line that appears as a borderline between the light blue (safe) and the white (dangerous) zones and clearly separates the two groups of subjects with different outcomes. The median fALFF values for G1 subjects are shown by a dashed line. [Color figure can be viewed at wileyonlinelibrary.com]
subjects receiving IMRT. There was a significant correlation between the PCC-to-right-hippocampus FC and the maximum dose applied to the right temporal lobe ($r = -0.66$, $P < 0.05$, see Fig. 7B). The higher the maximum dose, the lower the FC strength observed.

**DISCUSSION**

**General Discussion**

For the first time, we explored brain-wide RT-induced functional and structural abnormalities in normal-appearing nasopharyngeal carcinoma patients using the voxel-based analysis. We found that the brains without any visible lesions still had significant functional alterations at different post-RT stages (Figs. 2 and 3), particularly in the bilateral hippocampus. The functional changes seemed to be more sensitive than the white matter FA changes to the RT-induced injury, with the latter found to be in the splenium of the corpus callosum only after 12 months following the completion of RT, outside the zone traditionally thought to be irradiated (Fig. 4). We thus propose that the post-RT, normal-appearing brain was generally undergoing functional recovery with a structurally progressive course (Figs. 5 and 6). Interestingly, we found a dose-dependent PCC-to-right-hippocampus FC reduction during 6–12 months after RT (Fig. 7A), indicating that the DMN FC is quite sensitive to the...
irradiation. This suggests a careful choice of the RT dose, especially the dose to the right temporal lobe, is of great importance for the prevention of late-delayed neurodegeneration. Another clinically important finding is that the early increase in fALFF can be used as a predictor of temporal lobe necrosis that may occur many years later (Fig. 5B). These findings increase our understanding of post-RT brain injury, and emphasize the need to use rs-fMRI for monitoring post-RT brain function and tailoring RT strategy.

RT-induced brain damage is not a minor issue [Greene-Schloesser and Robbins, 2012; Tsai et al., 2014]. Most patients develop cognitive and psychiatric disorders in the late-delayed period that last for years [Cheung et al., 2003; Dietrich et al., 2008; Greene-Schloesser et al., 2013; Hsiao et al., 2010; Khong et al., 2006; Mo et al., 2014; Tang et al., 2012; Warrington et al., 2013; Wu et al., 2014]. Cognitive decline can occur in patients who have no MR-visible lesions [Chapman et al., 2012; Crossen et al., 1994], a finding that suggests the presence of microscopic lesions [Atwood et al., 2007; Matulewicz et al., 2006; Nagesh et al., 2008; Warrington et al., 2013; Ye et al., 2012]. Although much progress has been made [Abayomi, 1996; New, 2001], until now no consensus has been reached for human subjects on this question [Cheung et al., 2000]. We enrolled nasopharyngeal carcinoma patients to overcome the main constraints of previous studies for better study of general RT-induced injury, especially its presymptomatic stage (MR-invisible lesions), which is most meaningful for preventive care [Chan et al., 2004]. Our results also provide guidance for tailoring the dose and identifying the region to spare during radiotherapy [Gondi et al., 2010; Kazda et al., 2014]. Importantly, two models, a functional recovery while structural progression model and a preventive model, were proposed for future RT-injury studies (Fig. 5A) and were validated by follow-ups and retrospective observations (Fig. 5B). Our result, at least from the brain structure viewpoint, confirmed that the brain structure changes could be progressive, which is consistent with previous studies which found later increased FA abnormality [Chapman et al., 2012; Nagesh et al., 2008].

This is the first-reported fMRI-based RT-injury study. Several pioneering studies using other functional modalities—for example, PET, perfusion MRI, single-photon emission computed tomography (SPECT), and MRS—have been conducted mainly to investigate RT-induced necrosis [Chan et al., 2009b; Chen et al., 2014; Chong et al., 2001; Kim et al., 2010; Matulewicz et al., 2006; Schwartz et al., 1991; Sundgren, 2009; Tsai et al., 2000, 2001] or other MR-invisible lesions [Wang et al., 2012; Xiong et al., 2013]. These results can support our findings, as fMRI is also related to cerebral vascular function and perfusion. With rs-fMRI, more detailed spatiotemporal patterns of brain function can be investigated in vivo [Biswal, 2012; Raichle, 2010] due to its higher spatial and temporal resolutions, which provides new information that contributes to our understanding of RT injury obtained from the previous animal studies. Our results indicate that post-RT brain functional studies are of equal importance to white-matter structural studies because, although patients often recover from the observed functional abnormalities, these abnormalities may serve as early biomarkers of the severe temporal lobe necrosis that may develop many years later.

**Early Indicators of RT Injury**

One of our most important findings is the fALFF alterations in the early stage (0–6 months). Alterations at this stage have also been reported in previous studies using MRS. In a similar region, we previously found decreased N-acetylaspartic acid/choline and N-acetylaspartic acid/creatinine levels at 0-6 months after RT in another nasopharyngeal carcinoma cohort [Wang et al., 2012], particularly during the first 3 months after RT [Xiong et al., 2013]. Early decreases in metabolite levels have also been reported. For example, Chen et al. [2014] found that choline/creatine decreased to its lowest level at 3 months after RT. Sundgren et al. [2009] found a decreased metabolite ratio during RT, and the reduction lasted for 6 months. However, other studies have reported different results. For example, Chan et al. [2009b] found an RT-induced increase in choline/creatine in normal-appearing mouse brain, and Matulewicz et al. [2006] found a periodic fluctuation in choline/creatine during a long post-RT period in glioma patients. The inhomogeneity of the subject cohorts, the variety of irradiation targets, and the influence of surgery and tumor recurrence could contribute to such discrepancies. In this study, these nuisance effects were largely reduced.

The increased fALFF at the inferior temporal cortex in the early stage could reflect direct disruptions of cerebral vascular function, such as disruption of the blood–brain barrier due to vascular endothelial damage [Sundgren, 2009], vascular dilation [Wang et al., 2009], and increased vascularity [Schultheiss, 2012]. These changes may cause increased cerebral blood flow/volume, further causing hyperactivity. This explanation is further supported by studies involving perfusion MRI [Acker et al., 1998; Tsui et al., 2000, 2001]. In contrast, we think that neuronal dysfunction and demyelination [Chan et al., 2009b; Sundgren, 2009] are insignificant at the early stages because no decreased fALFF was found.

fALFF can be a radiosensitive early indicator. Focusing on the entire temporal lobe, we found a dynamic fALFF pattern that first increased, then decreased, and finally recovered (Fig. 2A–C). Based on the close relationship between fALFF and FC (Fig. 7A) and the common substrate between FC and white-matter integrity, we speculate that the early fALFF increase and the later fALFF/FC/FA decrease could be linked. Another important finding is that for the ten subjects in G2 whose outcome information was available, the fALFF increment in the inferior
temporal lobe at 0–6 months served as a predictive biomarker or early sign of severe late-delayed brain injury such as temporal lobe necrosis. That is, the early fALFF alterations could be regarded as an early sign of RT injury in the presymptomatic stage many years before clear MRI-based clinical indications were present. As indicated in Figure 5B, we proposed that if a subject has significantly increased fALFF in the lower part of the temporal lobe shortly after RT, this subject may have a higher probability of developing severe RT injury after years of progression. In contrast, if a subject has stable post-RT fALFF at its pre-RT level, he/she may be much more stable and show slower progressive brain changes. In Figure 5B, we tentatively used a threshold from the median fALFF value for G2 subjects, that is, $z = -0.44$, as a warning line. For patients with fALFF above this line, neuroprotective treatments to protect brain tissue from further damage are highly recommended. Owing to the high incidence of subject dropout, only 10 subjects in G2 were used for retrospective inspections. We are currently conducting a new retrospective study with a larger sample size to further validate the predictive value of fALFF in the presymptomatic stage of RT injury. As shown in Figure 5A, we have proposed a preventive model: by monitoring brain function in the early stages with fALFF from rs-fMRI, we could reduce or prevent fALFF abnormalities in the temporal lobe by means of early intervention; later reductions in the FC and white matter integrity and even temporal lobe necrosis could be ameliorated.

### Long-Term fALFF Fluctuations

In addition to the “increase-decrease-recover” pattern, we found that in specific areas of the inferior temporal lobe fALFF manifested dynamic changes with an “increase-decrease-increase” pattern (Fig. 2D, first column). This interesting fluctuation pattern is consistent with a previous report. Specifically, the interval between the two fALFF hyperintensities was approximately 6 months in our study, similar to an 8-month interval for the MRS metrics’ fluctuation period [Matulewicz et al., 2006]. Our result supports the hypothesis that this fluctuation is related to the dynamic interaction between blood-brain barrier disruption and brain repair processes [Matulewicz et al., 2006]. This finding also indicates that a neuroprotective mechanism might take effect after direct injury [Chen et al., 2014], which may be one source of brain functional recovery (the other source may be the rewiring of the brain connections, see “Temporarily Disrupted and Plastic FC in the DMN”).

### Late-Delayed fALFF Reductions

We found a late-delayed, extensive fALFF reduction in the right temporal cortex (Fig. 2B) at 6–12 months and scattered bottom-brain fALFF reductions even after 12 months (Fig. 2C). This long-lasting reduction is supported by previous studies [Sundgren, 2009] and could be due to brain volume loss or cumulative neurotoxic effects. Interestingly, we found that the reductions occurred in the gray matter near the white matter (Fig. 2B), which further supports the hypothesis that near-end white-matter neuronal demyelination or degradation is dominant in this midterm period (in contrast to earlier gray-matter abnormalities and later far-end callosal white-matter degradation). In addition to the near-end white-matter injury, gray-matter neuronal cell death caused by apoptosis and neuronal dysfunction secondary to irradiation might also have occurred [Chan et al., 1999].

In summary, we favor a dynamic multifactorial model of RT-induced local brain function injury. That is, in earlier periods, gray matter vasculature damage dominates RT injury, whereas later demyelination (from near-end to far-end white matter, which will be discussed later) dominates the RT aftereffects. Instead of supporting either a glial hypothesis [Hopewell and van der Kogel, 1999] or a vascular hypothesis [Lyubimova and Hopewell, 2004] of white-matter damage, we believe that both processes exist after RT but that the two processes dominate during different stages.

An interesting lateralization in RT injury was found. For examples, late-delayed fALFF reductions occurred mainly in the right temporal lobe, and dose-dependent FC changes were also concentrated in the right hippocampus (Fig. 7B). Currently, we have no clue to the cause of this asymmetric RT influence, but previous studies have reported similar focal degenerations in the same side of the temporal lobe [Chan et al., 2009a].

The inferior and superior frontal gyri displayed increased fALFF, and this increase became more significant over time (Fig. 2). Because the frontal area is outside the irradiation field, we speculate that this was due to compensatory effects in response to temporal lobe dysfunction. Considering that all these hyperactivities are in areas related to higher cognitive functions, this compensation could help to maintain normal cognitive function. Some poststroke longitudinal studies [Levin et al., 2009] and mild cognitive impairment studies [Qi et al., 2010] have also demonstrated such compensatory effects.

### Temporarily Disrupted and Plastic FC in the DMN

We found notable, distributed FC reductions in many DMN regions at 6–12 months after RT (Fig. 3). This is a significant “disruption and recovery” pattern which may indicate that the brain has the ability to recover. To our knowledge, this is a quite novel finding. This finding supports the cognitive decline, psychological disorders, and mood disorders commonly found after RT [Cheung et al., 2003; Greene-Schloesser et al., 2012; Khong et al., 2006; Mo et al., 2014; Tang et al., 2012; Wu et al., 2014] because the
DMN is essential for normal cognition and emotion [Buckner, 2013].

The decreased FC in the bilateral hippocampus is interesting. The hippocampus is an important node of the DMN [Buckner et al., 2008; Greicius et al., 2009] and is essential for learning and memory [Greicius et al., 2004]. The hippocampal cortex has dense reciprocal projections to the thalamus, midbrain, limbic system, and temporal lobe [Qin et al., 2015]. RT could impair these pathways and cause cognitive/affective disorders [Rocca et al., 2015] or directly impair the hippocampus itself [Son et al., 2015] by reducing hippocampal neurogenesis/proliferation [Conner et al., 2010; Schnegg et al., 2013] even at low radiation doses [Monje and Palmer, 2003]. Based on the dose-dependence of the FC in the hippocampus, we suggest sparing the hippocampus during RT [Gondi et al., 2010; Kazda et al., 2014] for better neurocognitive protection.

The reduction in the PCC-to-hippocampus FC is indicative of a similar phenomenon in patients with Alzheimer’s disease [Greicius et al., 2004], who also suffer from mild cognitive impairment in the preclinical stage. Akiyama et al. [2001] proposed a similar hypothesis that late-delayed RT injury to the white matter occurs mainly due to demyelination, similar to clinically accelerated brain aging. The difference between the plastic FC changes observed in our study and the continuously worsening FC in Alzheimer’s disease is related to the fact that our subjects had mild RT injury (MR-invisible lesions). In this cohort, the FC reduction in the late-delayed period was more flexible, possibly due to gradual cerebral vascular repair and increased numbers of neuronal cells in the affected regions [Chen et al., 2014]. Moreover, the increased FC in the frontal areas might act as a compensatory mechanism (a potential source of brain recovery following the lesions) that maintains normal cognitive function.

**Far-End White-Matter Progressive Damage**

Figure 4B shows clear, gradual FA reductions in the major white-matter structures, a novel and intriguing finding. DTI is a noninvasive technique that can detect white-matter damage in the absence of MR-visible lesions [Nagesh et al., 2008]. Some previous DTI studies have supported our finding of gradual white-matter damage [Nagesh et al., 2008; Welzel et al., 2008], whereas other studies have found that DTI metrics decreased in the early stage but partially recovered later [Chen et al., 2015; Wang et al., 2012; Xiong et al., 2013]. Animal studies [Chen et al., 2009b; Wang et al., 2009], however, support our finding by showing a progressive reduction in the FA with demyelination and a steeper decline when the RT dose was high. In addition, we think that the occurrence of abnormal regional functioning and inter-regional FC earlier (<12 months) than the white matter damage (>12 months) was reasonable because brain function might be more vulnerable or sensitive to attack [Karim et al., 2016]. Interestingly, similar white-matter regions (the periventricular areas) were also found to be frequently targeted in normal aging subjects [Habes et al., 2016]. Taken together, we think our results of the white-matter progressive damage are reliable.

One possible reason for the lack of earlier stage FA reduction is that, as previously discussed, the partial volume effect and the possible registration error, particularly in the gray and white matter interface, may prevent DTI from sensitively detecting subtle changes [Schultheiss, 2012]. Therefore, our results did not indicate intact white matter in the earlier stages; rather, the white matter lesions became larger and more prominent later on [Chapman et al., 2012] and finally caused significant reductions in FA. Further studies should focus on this subtle white matter injury in the acute response stage after RT using more advanced diffusion imaging techniques with a higher spatial resolution because this earlier response could predict late injury [Chapman et al., 2012; Chen et al., 2015]. Another more likely reason for the lack of earlier stage FA reduction is that the different analytical methods used in our study (voxel-wise comparison) compared to several previous studies (ROI analysis) [Chen et al., 2015; Wang et al., 2012; Xiong et al., 2013] produced different results. In the ROI-based studies, only a small area was selected from the entire inferior longitudinal fasciculus in each subject; there is no further constraint for the ROI placement. Human intervention in the relatively subjective ROI selection has posed a serious problem. A small change in the location of the ROI may lead to large variations in FA values and biased group comparison results. Therefore, these ROI-based results may have a reliability issue. Our voxel-wise whole-brain analysis is more objective and thus more reliable.

**Links Between Functional and Structural Changes**

An important finding of this study is that, for the first time, we were able to link earlier RT-induced functional abnormalities with later impaired white matter integrity using DTI tractography (Fig. 6). This is also the basis of our proposed “preventive model” (Fig. 5A). Putative neuronal fiber bundles were found that connect the functional changes in the gray matter of the temporal lobe to the structural changes in the white matter outside the temporal lobe. This finding has also received support from RT-induced late response studies in which the periventricular white matter was found to undergo detectable change only after 12 months [Constine et al., 1988; Packer et al., 1986] and the large fiber bundles in the genu and splenium of the corpus callosum were found to undergo progressive structural degradation after one year [Nagesh et al., 2008], which correlated with impaired cognition [Akiyama et al., 2001]. Based on our preventive model,
Our results suggest that (1) as a promising early sign, increased local brain activity in the inferior temporal lobe could be used to monitor early RT-induced brain injury at the presymptomatic stage of RT injury and to predict the occurrence of severe brain necrosis many years later; (2) the FC between the bilateral hippocampus and the PCC could be the cause of a broad spectrum of post-RT cognitive impairments as late responses; (3) whereas the FC disruption is plastic and reversible in normal-appearing brains, the potentially degenerative white-matter damage lasts longer, could be progressive, and could be a substrate of late toxicity and long-term cognitive abnormality; (4) the RT dose, particularly the maximum dose at the temporal lobe, should be carefully determined, and the course of RT should be closely monitored using fMRI; and (5) the practitioner should consider sparing the hippocampus or even the entire temporal lobe during RT.

Limitations

This study has several limitations. First, our sample size was relatively small because we used stringent subject inclusion criteria; the subjects with distant metastases were excluded to avoid any protopathic brain invasion that could interfere our results. We also only included the patients in the presymptomatic stage to explore the potential of early detection and to detect early biomarkers for developing future preventive interventions. The patients with this type of lesions are far less than those with visible lesions. In addition, we further calculated the effect size of our metrics and found that most of our results had small effect sizes. In addition, the intervention should be applied not only to the temporal lobe but also to the fiber bundles that are at the far end and have dense connections to the traditionally expected irradiation damaging zone.

Clinical Implications

Using precious multimodal brain imaging data, we revealed how the postradiotherapy brain undergoes an evolutional process with both functional recovery and structurally degenerative progression during the presymptomatic stage of RT injury. We have proposed that dynamic multifactorial processes occur in the irradiation injured presymptomatic brain involving earlier vasculopathy dominance and later demyelination dominance. Our results suggest that both progressive white matter degeneration from the near- to the far-end and neuroprotective processes and compensatory effects are involved in this recovery/progression model. Our additional preventive model highlights the importance of early brain function monitoring, careful RT-dose tailoring and timely preventive neuroprotection for avoiding irreversible severe complications that worsen outcomes.

CONCLUSION

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