The Temporal Dynamics of Brain Plasticity in Aging

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Abstract

Cognitive training has been suggested as a possible remediation of decline in brain structure with older age. However, it is unknown whether training effects are transient or enduring, as no studies have examined training-induced plasticity relative to decline in older adults across extended periods with multiple intervention phases. We investigated the temporal dynamics of brain plasticity across periods on and off memory training, hypothesizing that (1) a decline in white matter (WM) microstructure would be observed across the duration of the study and (2) that periods of memory training would moderate the WM microstructural decline. In total, 107 older adults followed a 40-week program, including 2 training periods separated by periods with no intervention. The general decline in WM microstructure observed across the duration of the study was moderated following the training periods, demonstrating that cognitive training may mitigate age-related brain deterioration. The training-related improvements were estimated to subside over time, indicating that continuous training may be a premise for the enduring attenuation of neural decline. Memory improvements were largely maintained after the initial training period, and may thus not rely on continuous training to the same degree as WM microstructure.

Key words: aging, cognitive training, memory, plasticity, white matter microstructure

Introduction

Experience-dependent brain plasticity is well documented in both young and older adults (Draganski et al. 2004; Boyke et al. 2008; Scholz et al. 2009). Targeted cognitive training has thus been suggested as a possible remediation of decline in brain structure with older age. Although the evidence for a lifelong potential for plasticity provides a promising outlook, the premises for, and limitations of, plasticity is far from fully understood. Studies of young adults indicate that neural alterations do not persist in the absence of exercise (Draganski et al. 2004), and that structural changes recede during subsequent training on the same skill (Driemeyer et al. 2008). Thus, training effects may not last beyond the intervention periods, and the learning of new skills may be more effective for the mitigation of decline relative to sustained training on the same tasks. In older adults, brain aging itself is likely to influence the time course of plasticity, such that training effects need to be measured in relation to general decline over time. However, no studies have examined training-induced plasticity relative to decline in older adults across extended periods with multiple intervention phases. Hence, it is unknown whether repeated cognitive interventions can systematically moderate the magnitude of brain deterioration in aging, and whether continuous training is a premise for the attenuation of neural decline. White matter (WM) microstructure is highly susceptible to age-related deterioration (Sexton et al. 2014; Bender et al. 2016), and short-term changes in WM have been observed across periods of less than 3 months in healthy older adults (Engvig et al. 2012). Can WM trajectories be modified, and if so, by what means?
Evidence suggests that both young and older adults can benefit from memory strategy training (Cavallini et al. 2003; Engvig et al. 2012; de Lange et al. 2017). As this type of training has been shown to influence WM microstructure in older adults (Engvig et al. 2012; de Lange et al. 2017), we investigated whether periods of memory strategy training could modify trajectories of WM decline in 107 older adults (mean age ± SD = 73.2 ± 2.9), by using an ABAB/BABA design as illustrated in Figure 1. The training aimed to improve serial verbal recollection memory by implementing the mnemonic technique Method of Loci (MoL). An active control group focusing on popular science was included during the first 10-week period to investigate the specificity of the memory-training effects. All participants were examined with magnetic resonance imaging (MRI) and cognitive testing, with a 10-week interval between each assessment. Microstructural changes were measured using the diffusion tensor imaging (DTI) derived metrics fractional anisotropy (FA), mean diffusivity (MD), radial diffusivity (RD), and axial diffusivity (AD). We hypothesized that (1) a decline in WM microstructure would be observed over 40 weeks, (2) memory training would moderate the WM microstructural decline, and (3) training effects on WM microstructure would subside in the absence of training, but (4) would be reinvoked with a new training period.

Materials and Methods

Sample

The sample was drawn from the project “Neurocognitive Plasticity” at the Center for Lifespan Changes in Brain and Cognition (LCBC), Department of Psychology, University of Oslo. All procedures were approved by the Regional Ethical Committee of Southern Norway, and written consent was obtained from all participants. Participants were recruited through newspaper and webpage adverts, and were screened with a health interview. Participants were required to be between 70 and 80 years old, healthy adults, right-handed, fluent Norwegian speakers, and have normal or corrected to normal vision and hearing. Exclusion criteria were history of injury or disease known to affect central nervous system (CNS) function, including neurological or psychiatric illness or serious head trauma, being under psychiatric treatment, use of psychoactive drugs known to affect CNS functioning, and MRI contraindications. All scans were evaluated by a neuroradiologist and deemed to be free of significant injuries or conditions. For inclusion in the study, participants were required to score ≥ 26 on the Mini Mental State Examination (MMSE) (Folstein et al. 1975) and have scores less than 2 standard deviations below mean on the 5 min delayed recall subtest of the California Verbal Learning Test II (CVLT II) (Delis et al. 2000). Three individuals were excluded based on these criteria. All participants further had to achieve an IQ above 85 on the Wechsler Abbreviated Scale of Intelligence (WASI) (Wechsler 1999). A total of 125 participants fulfilled the inclusion criteria. Sample demographics are shown in Table 1.

Design and Training Intervention

Pools of around 20 participants were recruited at a time, and the participants were assigned to 1 of 3 intervention groups at registration. The data collection was on-going and continuous for the 3 conditions simultaneously, ensuring that participants from all 3 experimental groups were scanned and tested interchangeably and thus reducing the possibility of group differences with regard to the assessment and scanning conditions. Although group assignments based on date do not comply with suggested criteria for randomization of participants (Schulz and Grimes, 2002), practical considerations forced a compromise due to the extensive data collection with strict time intervals and assessments locked to specific dates across 40 weeks. Group 1 (ABAB) started with 10 weeks of memory training (N = 57), and moved on to a subsequent rest period with no intervention followed by a second training period, and then a final rest period. Group 2 (BABA) started with 10 weeks of rest (N = 50), and moved on to the first training period followed by a rest period and a subsequent training period. The participants allocated to groups 1 and 2 completed scanning and cognitive testing at 5 occasions, with a 10-week interval between the assessment sessions. An active control group (group 3, AC, N = 18) was included during the first period in order to investigate the specific cognitive effects of the memory training. The active control participants completed scanning and cognitive testing at 2 occasions, with a 10-week interval between the assessment sessions.

The memory-training program aimed at improving serial verbal recollection memory by implementing the mnemonic technique MoL (Bower 1970), which has been shown to improve serial recall substantially in older adults (Engvig et al. 2010; de Lange et al. 2016). The training program included a single course session each week and home assignments involving memorizing word lists. The first group session included a presentation of the project, an introduction to the MoL with instructions, and an initial word list task consisting of 15 words. The following weekly group sessions included updating of the strategy, clarification of instructions and a word list task, which was increased by 5 words each week to ensure a continuous challenge. However, the participants were encouraged to individually adjust the difficulty level, with the aim of achieving a challenging but manageable training level across all the participants. Individual adjustment involved increasing/decreasing the number of words on the tasks to a sufficiently challenging level, performing the tasks within individual time limits and recollection of the word lists in reverse order. Although the exact number of words was subject to individual adjustment, all participants completed the training with a weekly increase in number of words. Eight home assignments were sent out weekly, with a minimum requirement that 4 be

![Figure 1](https://example.com/figure1.png)

Figure 1. The course of the training program depicted for each of the 3 groups. In total, 71 participants completed the full study. Drop-out rates are provided in the Supplementary Material.
completed. The home assignments consisted of word lists with various themes and followed the level of difficulty set in the group session the same week. The tasks included options for individual adjustment including increasing/decreasing the number of words, performing the tasks within individual time limits and recollection of the word lists in reverse order. The home assignments were completed online. All responses in addition to time spent on the tasks were registered to a database. The proportion of tasks completed across participants was 74% during the first training period, and 69% during the second period. The 2 training periods followed the same structure and included the same strategy training. The initial level of difficulty in the second training period corresponded to the middle level of the first period. The weekly increase in number of words in the lists was doubled during the last 3 weeks of the second training period. Thus, the final level of difficulty was higher in the second training period. The participants were instructed not to practice any memory training in the rest periods. The active control group program involved popular scientific lectures once a week. Eight home assignments were sent out weekly. The home assignments were completed online, and involved tasks related to the weekly popular scientific themes. The proportion of tasks completed across participants in the active control group was 70%. None of the tasks or lectures in the active control program involved any specific form of memory training. Contact with staff, group meetings and the number of tasks were matched between the training group and the active control group. Independent sample t-tests (2-sided) showed that the number of tasks completed did not differ between the training group and the active control group (mean ± SD = 57.6 ± 15.1 [range = 76] for the training group, and 55.5 ± 21.1 [range = 72] for the active control group, t(60) = -0.45 P = 0.7). We have recently investigated the influence of number of tasks completed on memory improvement and WM microstructure in the same sample. Number of tasks did not influence memory improvement or WM microstructural change during the first 10-week period (de Lange et al. 2017). Test sessions and time intervals were held identical for all participants, in order to ensure that test-retest effects would not differ across the groups.

Image Acquisition and Data Processing

A Siemens Skyra 3T MRI scanner with a 24-channel head-coil was used (Siemens Medical Solutions; Erlangen, Germany). A diffusion-weighted echo-planar imaging (EPI) sequence was applied for each subject (FOVxy = 252 × 256 mm, dimensions = 128 × 130 × 70, voxel size = 1.9626 × 1.9626 mm³, slice thickness = 2 mm, repetition time = 9300 ms, echo time = 87 ms). Overall, 64 unique diffusion-weighted volumes were collected at b-value = 1000 s mm⁻¹ in addition to 2 nondiffusion-weighted (b-value = 0 s mm⁻¹) volumes, one acquired with an opposite k-space traversal direction for the purpose of correcting susceptibility artefacts. All scan-sets were manually checked for gross motion artefacts. The susceptibility-induced field was estimated using the FSL tool topup (Andersson et al. 2003) and corrected for along with subject motion and eddy current-induced fields using the eddy tool (Andersson and Sotiropoulos 2016). Signal dropout caused by subject motion during the diffusion encoding was detected and corrected using the eddy tool, as implemented in FSL (Andersson and Sotiropoulos 2016). Each acquired slice was compared with a model free prediction, and if the observed signal was statistically different (3 standard deviations) from the prediction, it was replaced by the latter. An average of 0.38, 0.38, and 0.40 slices per volume across subjects were replaced in groups 1, 2, and 3, respectively. The number of slices replaced did not differ between groups (one-way analysis of variance (Bonferroni corrected) showed no differences between the groups on the above measures.

### Statistical Analyses

#### The Time Course of WM Microstructural Changes

For each subject at every time point, the average MD, RD, AD, and FA values were calculated within the mean FA skeleton mask. The time course of WM microstructural changes was analyzed using a nonlinear mixed effects model, using the default covariance pattern (http://uk.mathworks.com/help/...
Memory improvement was measured using an experimental word-list test developed to measure verbal recollection. The test enabled the MoL to be applied, such that the measure of memory performance was closely related to the utilized technique, and thus convenient for measuring training gains (see de Lange et al. (2017) for details). The total number of words recalled from the word-list test was used as the measure of memory performance. In a similar fashion to microstructural changes, memory trajectories were analyzed using a nonlinear mixed effects (NLME) model, as implemented in Matlab. First, improvements relative to the intercept were estimated across the duration of the study, using the following function:

\[ \text{Memory}(t) = \beta_0 + \beta_1 \times e^{-\frac{t}{\beta_2}} + \beta_3 \times e^{-\frac{t}{\beta_4}} + \beta_5 \times \text{age} + \beta_6 \times \text{sex} \]

The terms \( \beta_1 \) and \( \beta_3 \) model changes in memory performance relative to the intercept (\( \beta_0 \)), while \( \beta_2 \) and \( \beta_4 \) model the times at which the changes occur. The intercept (\( \beta_0 \)) and improvement phases (\( \beta_2 \) and \( \beta_4 \)) were modeled as random factors. The remaining beta terms model the fixed effects of age and sex across all time points. Next, we calculated the difference between the estimated performance at the time points of improvement and the estimated performance before each improvement phase.

To assess possible transfer effects during the first 10-week period, repeated measure ANCOVAs were run using intervention group as between-subject factor, and change in performance on the following tests as dependent variables: The Wechsler Digit Span test, a working memory test where the participants were asked to render a sequence of numbers forward and backward, the Rey–Osterrieth Complex figure test (RCFT), measuring visual recollection (Lezak 2004), an experimentally developed paired-words test, where the participants were asked to recall pairs of unassociated words, and the CVLT learning and recall trials. The memory training did not influence performance on the Digit Span test, the RCFT or the paired-words test. However, performance on the CVLT learning and 30-min recall improved to a larger extent in the training group relative to the control group, reflecting a near-transfer effect. The results are provided in the Supplementary Material.

Results

The Time Course of Microstructural Changes

The time course of MD is shown in Figure 2. The slope function (\( \beta_0 \)) estimated a positive slope, indicating a general increase in MD across the duration of the study. An improvement in MD
(decreased MD relative to the slope, $\beta_3$) was estimated to begin during the first training period (between time points 1 and 2), and to wear off during the subsequent rest period (between time points 2 and 3). The model further found a second improvement in MD ($\beta_4$), which was estimated to begin during the second training period (as shown by the turning point in the curve between time points 3 and 4), followed by a return towards the slope in the final rest period (between time points 4 and 5). A summary of the results is presented in Table 2.

The time courses of AD, RD, and FA are shown in Figure 3. The time courses of AD and RD showed similar results to those of MD, with a positive slope, indicating an increase in these metrics across the duration of the study. An improvement in AD and RD (decreased AD and RD relative to the slope, $\beta_3$) was estimated to begin during the first training period (between time points 1 and 2), and to wear off during the subsequent rest period (between time points 2 and 3). The model further found a second improvement in AD and RD ($\beta_4$), which was estimated to begin during the second training period (as shown by the turning point in the curve between time points 3 and 4), followed by a return towards the slope in the final rest period (between time points 4 and 5). The time course of FA showed variations corresponding to the opposite of those of MD, AD, and RD. The slope function demonstrated a negative slope, indicating a general decrease in FA across the duration of the study. An improvement in FA (increased FA relative to the slope, $\beta_2$) was estimated to begin during the first training period (between time points 1 and 2), and to wear off during the subsequent rest period (between time points 2 and 3). The model further found a second improvement in FA ($\beta_4$), which was estimated to begin during the second training period (as shown by the turning point in the curve between time points 3 and 4), followed by a return towards the slope in the final rest period (between time points 4 and 5).

To compare the magnitude of WM microstructural changes in the 2 improvement phases, we first calculated the mean differences in MD and FA during each of the phases estimated by the model, and then calculated their z-score difference using the following equation:

$$z = \frac{(\delta_1 - \delta_2)}{\sqrt{SE_{\delta_1}^2 + SE_{\delta_2}^2}}.$$

The parameters $\delta_1$ and $\delta_2$ represent the magnitude of change in the model during each improvement phase. SE represents the standard errors. The 2 improvement phases did not differ for MD ($\delta_1 \pm SE = (0.93 \pm 4.05) \times 10^{-6}$, $\delta_2 = (7.24 \pm 8.79) \times 10^{-6}$, $z = -0.46$, $P = 0.64$) or FA ($\delta_1 = (-0.28 \pm 1.94) \times 10^{-3}$, $\delta_2 = (2.00 \pm 3.75) \times 10^{-3}$, $z = 0.30$, $P = 0.77$).

To investigate effects in specific tracts, the model was run on MD in the inferior frontal–occipital fasciculus (IFOF), inferior longitudinal fasciculus (ILF), superior longitudinal fasciculus (SLF), uncinate fasciculus (UF), and the hippocampal cingulum bundle (HCB), based on the JHU DTI-based WM atlas ( Mori et al. 2005). Bonferroni’s adjustment was applied to correct for multiple comparisons. The slope function demonstrated a positive slope for all the tracts (ILF: $\beta_1 = [7.00 \pm 1.90] \times 10^{-6}$, $F[1, 419] = 13.60$, $P = 2.56 \times 10^{-4}$, SLF: $\beta_1 = [5.49 \pm 1.69] \times 10^{-6}$, $F[1, 419] = 10.49$, $P = 0.001$, HCB: $\beta_1 = [1.42 \pm 0.39] \times 10^{-5}$, $F[1, 419] = 13.06$, $P = 3.38 \times 10^{-4}$, UF: $\beta_1 = [6.08 \pm 2.47] \times 10^{-6}$, $F[1, 419] = 6.08$, $P = 0.01$, ifOF: $\beta_1 = [4.98 \pm 1.80] \times 10^{-6}$, $F[1, 419] = 7.64$, $P = 0.01$), indicating a general increase in MD across the duration of the study. An improvement in MD (decreased MD relative to the slope, $\beta_3$) was found in ILOF ($\beta_3 = [-1.23 \pm 0.43] \times 10^{-5}$, $F[1, 419] = 8.41$, $P = 0.003$) and HCB ($\beta_3 = [-2.75 \pm 0.90] \times 10^{-5}$, $F[1, 419] = 9.20$, $P = 0.003$). This improvement was estimated to begin during the first training period (between time points 1 and 2), and to wear off during the subsequent rest period (between time points 2 and 3). In these tracts, the model further found a second improvement phase ($\beta_4$) in SLF: $\beta_4 = [-2.85 \pm 0.95] \times 10^{-5}$, $F[1, 419] = 9.12$, $P = 0.003$, HCB: $\beta_4 = [-6.34 \pm 1.20] \times 10^{-5}$, $F[1, 419] = 10.51$, $P = 0.001$), which was estimated to begin during the second training period (as shown by the turning point in the curve between time points 3 and 4), followed by a return towards the slope in the final rest period (between time points 4 and 5). No significant MD improvements ($\alpha$ level for Bonferroni-corrected P-values = 0.01) were found for SLF ($\beta_3 = [-8.71 \pm 3.88] \times 10^{-6}$, $F[1, 419] = 5.03$, $P = 0.03$, $\beta_4 = [-1.71 \pm 0.84] \times 10^{-5}$, $F[1, 419] = 4.12$, $P = 0.04$), ifOF: $\beta_3 = [-6.85 \pm 4.16] \times 10^{-6}$, $F[1, 419] = 2.71$, $P = 0.10$, $\beta_4 = [-1.65 \pm 0.87] \times 10^{-5}$, $F[1, 419] = 3.62$, $P = 0.06$ and UF ($\beta_3 = [-9.09 \pm 5.89] \times 10^{-6}$, $F[1, 419] = 2.38$, $P = 0.12$, $\beta_4 = [-1.97 \pm 1.21] \times 10^{-5}$, $F[1, 419] = 2.63$, $P = 0.12$). The time course of MD in SLF and HCB is shown in Figure 4. The spatial locations of the tracts are shown in Figure 5.

**Table 2** WM microstructure results.

| WM       | Parameter | Description                  | Value ± SE | F(1, 419) | P       |
|----------|-----------|------------------------------|------------|-----------|---------|
| MD       | $\beta_1$ | Slope                        | (6.21 ± 1.77) × 10^{-6} | 12.29 | 5.03 × 10^{-4} |
|          | $\beta_2$ | Improvement phase 1          | (-9.54 ± 4.05) × 10^{-6} | 5.54 | 0.02 |
|          | $\beta_3$ | Improvement phase 2          | (-21.70 ± 8.79) × 10^{-6} | 6.10 | 0.01 |
| AD       | $\beta_1$ | Slope                        | (6.63 ± 1.35) × 10^{-6} | 8.00 | 0.01 |
|          | $\beta_2$ | Improvement phase 1          | (-1.07 ± 0.52) × 10^{-5} | 4.19 | 0.04 |
|          | $\beta_3$ | Improvement phase 2          | (-2.34 ± 1.14) × 10^{-5} | 4.15 | 0.04 |
| RD       | $\beta_1$ | Slope                        | (5.96 ± 1.56) × 10^{-6} | 14.63 | 1.51 × 10^{-4} |
|          | $\beta_2$ | Improvement phase 1          | (-8.92 ± 3.61) × 10^{-6} | 6.09 | 0.01 |
|          | $\beta_3$ | Improvement phase 2          | (-2.11 ± 0.78) × 10^{-5} | 7.43 | 0.01 |
| FA       | $\beta_1$ | Slope                        | (-2.09 ± 0.79) × 10^{-3} | 6.97 | 0.01 |
|          | $\beta_2$ | Improvement phase 1          | (3.40 ± 1.94) × 10^{-3} | 3.07 | 0.08 |
|          | $\beta_3$ | Improvement phase 2          | (7.55 ± 3.75) × 10^{-3} | 4.06 | 0.05 |

Results of the nonlinear mixed effects model for each white matter metric in the full skeleton. The 2 improvement phases were estimated to begin during each training period. $\beta_3$ represents the degree of improvement relative to the slope in improvement phase 1. $\beta_4$ represents the degree of improvement relative to the slope in improvement phase 2. MD = mean diffusivity; AD = axial diffusivity; RD = radial diffusivity; FA = fractional anisotropy.

The Time Course of Memory Performance

The time course of memory performance is shown in Figure 6. The model estimated 2 phases of memory improvement.
relative to the intercept (β1 ± SE = 10.62 ± 2.24, F[1, 417] = 22.50, P = 2.89 × 10−6, β3 = 10.84 ± 4.51, F[1, 417] = 5.78, P = 0.02), which corresponded to the phases of WM microstructural improvements. Memory performance changed significantly during the first improvement phase (mean change ± SE = 8.50 ± 1.05, z = 5.71, P = 1.0 × 10−6), while the change in performance was not significant during the second improvement phase (mean change ± SE = 2.01 ± 1.33, z = 1.07, P = 0.28). A comparison of the 2 improvement phases (using the same equation as for MD and FA) showed a larger memory change during the first improvement phase relative to the second (z = 2.70, P = 0.007).

Correlations between WM microstructure and memory performance at each time point of assessment showed no relationship between MD and memory performance at time point 0 (r = 0.17, P = 0.30), while negative relationships were observed between MD and memory performance at time points 1 (r = −0.26, P = 0.01),
not forced to occur at the exact position of the measured time points of assessment. Hence, although the continuous function estimates a value at every point across the duration of the study, additional time points of assessment would have enabled a more precise description of the observed changes.

Although the physiological basis for water diffusion in WM is not fully understood (Beaulieu 2002), evidence suggests that WM microstructural decline is reflected by decreased FA and increased diffusivity, which is commonly observed in aging (Bennett and Madden 2014; Sexton et al. 2014). The memory training moderated the magnitude of the age-related decline in WM microstructure, corresponding to previous studies showing training-related microstructural plasticity in older adults (Lövdén, Bodammer, et al. 2010; Bennett et al. 2011; Engvig et al. 2012). DTI measurements reflect the restriction of water molecule motion, which can be imposed by cellular properties such as membranes, axonal density and myelin (Beaulieu 2002). While MD measures general diffusivity, AD and RD represent the rate of diffusion along the primary and secondary axes of the diffusion ellipsoid, respectively. FA measures the difference between the largest eigenvalue relative to the others and has been associated with restricted molecular motion caused by directionally oriented microstructures such as myelin sheaths and axonal cell membranes (Pierpaoli et al. 1996; Beaulieu 2002).

The observed trajectories of MD, AD, and RD changes were accompanied by changes in FA in the opposite direction. Decreased MD, AD, and RD was accompanied by increased FA in the mitigation periods, while the opposite pattern was observed for the general slope across the duration of the study. Some evidence indicates that age-related decrease in FA is primarily driven by an increase in RD (Rbhat and Beaulieu 2004; Madden et al. 2009). Given the associations between both FA and RD and myelin in animal studies (Song et al. 2005; Blumenfeld-Katzir et al. 2011), the alterations in these metrics could be driven by changes in myelination. Recent animal studies have shown training-related increases in immunofluorescence staining of myelin basic protein, which is indicative of myelination, in co-occurrence with increased FA (Blumenfeld-Katzir et al. 2011; Sampaio-Baptista et al. 2013). Myelination is also suggested to be a central for human learning (Fields 2008, 2010). Although myelination may modulate the degree of anisotropy, evidence suggests that axonal membranes largely contribute to anisotropic diffusion (Beaulieu 2002). Thus, the changes in FA may also have been affected by the condition of axonal membranes. While the observed variations in AD could be associated with axonal alterations (Song et al. 2003), and the coherence of axonal orientation (Bennett II, DJ Madden, CJ Vaidya, DV Howard, and JH Howard, Jr. 2010), the changes in MD could indicate underlying alterations in relatively isotropic structures such as astrocytes (Blumenfeld et al. 2006; Sagi et al. 2012), which has been observed in animals after training (Blumenfeld-Katzir et al. 2011; Sagi et al. 2012). Alternatively, reduced MD may be driven by myelination of axons in crossing fiber regions (Mackey et al. 2012). Thus, the mechanisms underlying changes in DTI metrics depends upon the local fiber architecture. Clearly, signal changes from DTI require careful interpretation, as the exact neurobiological underpinnings of diffusion metrics cannot be directly inferred. Although DTI may be sensitive to underlying cellular changes of large enough volumetric contribution (Sagi et al. 2012; Fields 2015), the signal is also influenced by how axons are laid out within the voxel, as the gradients are applied along given axes (Jones et al. 2013).

Diffusion measurements are also prone to cerebrospinal fluid...
(CSF) based partial volume artefacts (Alexander et al. 2001; Metzler-Baddeley et al. 2012). As aging is associated with white and grey matter loss, the present study could benefit from corrections for CSF contamination to improve the precision of the WM measures. However, it has been reported that the commonly observed age-related differences in DTI metrics cannot be attributed to partial volume effects (Pfefferbaum and Sullivan 2003; Bhagat and Beaulieu 2004). Although some aging studies have indicated region-specific patterns of increased RD and reduced AD (Bennett et al. 2010; Burzynska et al. 2010), the predominant picture appears to be one of increased diffusivity in general. Changes in DTI metrics have previously been observed across periods of less than 3 months in older adults (Engvig et al. 2012), indicating that diffusion MRI is sensitive to short-term changes in microstructure. The general decline in WM microstructure across the study corresponded to the estimated annual change in a matched non-intervention sample, suggesting that age-related decline in WM microstructure is detectable over a period of less than a year.

As the cognitive processes involved in mnemonic strategies are likely to rely on multiple brain areas, efficient transfer and integration of information between these distributed regions is critical. Although the overall evidence does not currently demonstrate a high degree of regional specificity in the relationship between WM integrity and cognition (Madden et al. 2009; Salthouse 2011; Dresler et al. 2017), the observed differences in the extracted tracts indicate that the inferior longitudinal fasciculi and the hippocampal cingulum bundle may represent tracts of particular importance for information transfer that is beneficial for cognitive improvements after memory strategy training.

As evidence suggests that learning a new skill may affect brain structure to a greater extent relative to practices those already learned (Driemeyer et al. 2008; Lövdén, Bäckman, et al. 2010), it is possible that a second training period comprising a new type of training could boost the neural mitigation to a larger extent. Thus, although beyond the scope of the present study, comparing the time course of plasticity in response to different training paradigms, for instance by also applying processed based working memory training (Karbach and Verhaeghen 2014), may represent a key focus for future research.

**Conclusion**

Targeted training has the potential to moderate the magnitude of age-related brain deterioration. The training-related improvements in WM microstructure were estimated to subside over time, indicating that continuous training may be a premise for the enduring attenuation of neural decline. Memory improvements from the initial training period were largely maintained across the duration of the study. Cognitive improvements may thus not rely on consistent training to the same degree as WM microstructure. In conclusion, the results suggest that continuous engagement in cognitive activities may influence brain-aging trajectories.

**Supplementary Material**

Supplementary material is available at Cerebral Cortex online.

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**Notes**

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