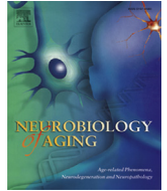




Contents lists available at ScienceDirect

Neurobiology of Aging

journal homepage: www.elsevier.com/locate/neuaging

Structural complexity is negatively associated with brain activity: a novel multimodal test of compensation theories of aging



Ian M. McDonough^{a,*}, Christopher R. Madan^b

^a Department of Psychology, The University of Alabama, Tuscaloosa, AL, USA

^b School of Psychology, University of Nottingham, Nottingham, UK

ARTICLE INFO

Article history:

Received 29 February 2020

Received in revised form 13 October 2020

Accepted 24 October 2020

Available online 14 November 2020

Keywords:

Aging

Compensation

Default mode network

Episodic memory

Fractal dimensionality

fMRI

ABSTRACT

Fractal dimensionality (FD) measures the complexity within the folds and ridges of cortical and subcortical structures. We tested the degree that FD might provide a new perspective on the atrophy-compensation hypothesis: age or disease-related atrophy causes a compensatory neural response in the form of increased brain activity in the prefrontal cortex to maintain cognition. Brain structural and functional data were collected from 63 middle-aged and older adults and 18 young-adult controls. Two distinct patterns of FD were found that separated cortical from subcortical structures. Subcortical FD was more strongly negatively correlated with age than cortical FD, and cortical FD was negatively associated with brain activity during memory retrieval in medial and lateral parietal cortices uniquely in middle-aged and older adults. Multivariate analyses revealed that the lower FD/higher brain activity pattern was associated with poorer cognition—patterns not present in young adults, consistent with compensation. Bayesian analyses provide further evidence against the modal interpretation of the atrophy-compensation hypothesis in the prefrontal cortex—a key principle found in some neurocognitive theories of aging.

© 2020 Elsevier Inc. All rights reserved.

1. Introduction

As the world's aging population grows, so too does the risk for cognitive decline and age-related neurodegenerative disorders such as Alzheimer's disease (AD). Understanding and preventing such declines in cognition is critical to increase the length of time that one leads a happy and independent life (Rowe and Kahn, 1987). Over the past 3 decades, neuroimaging has been used as a tool to understand how age differences in the brain might inform the causes and consequences of cognitive aging (for a brief review, see Park and McDonough, 2013). This literature has generally revealed declining brain structures (Fjell et al., 2014) alongside patterns of both higher and lower brain activity (for meta-analyses, see Li et al., 2015; Spreng et al., 2010). In the present study, we took a multimodal approach using a relatively novel measure of brain structure, fractal dimensionality (FD), to investigate the relationship between brain structure and function in the aging brain.

A prominent hypothesis arising from the past 15 years of neuroimaging research in aging is that brain atrophy, due to advanced aging or disease, causes an increased neural response in an attempt to maintain cognition (for review, see Cabeza et al., 2018). Although lower brain activity with aging also has been widely found, these patterns have been intuitively linked to a lack of brain maintenance (Nyberg et al., 2012), neural inefficiency (Logan et al., 2002), or decline in neural distinctiveness (Li et al., 2001). More counterintuitive and controversial is the notion of compensation in which atrophy in aging adults, largely in the prefrontal cortex (PFC), is offset by an increase in brain activity in nearby or contralateral PFC, and some research has extended this notion to the lateral parietal cortex (LPC) (Cabeza, 2002; Greenwood, 2007; Park and Reuter-Lorenz, 2009; Reuter-Lorenz and Cappell, 2008). The PFC often is implicated in executive control processes to flexibly adapt to ongoing task demands across many tasks (e.g., Vincent et al., 2008; Power and Petersen, 2013). One of the first studies to document higher age-related PFC activity was in an episodic memory task (e.g., Cabeza et al., 2002), and subsequent research investigating compensatory activity also has used episodic memory tasks during either encoding or retrieval (Brassen et al., 2009; Cabeza et al., 2002; Düzel et al., 2011; Persson et al., 2012; Pudas et al., 2013; Rajah et al., 2011). Supporting these single studies, quantitative meta-analyses have confirmed reliably higher brain activity in the

Role of funding source: Funding was provided by The University of Alabama through startup funds to I.M., the University of Alabama College Academy of Research, Scholarship, and Creative Activity to I.M., and the University of Alabama, Birmingham/The National Institutes of Health, grant/award number: P30AG031054.

* Corresponding author at: Department of Psychology, The University of Alabama, Tuscaloosa, AL 35487-0348, USA. Tel.: +1-205-348-1168; fax: +1-205-348-8648.

E-mail address: immcdonough@ua.edu (I.M. McDonough).

PFC in older relative to younger adults at memory encoding (Li et al., 2015; Spreng et al., 2010) and retrieval (Li et al., 2015; Spreng et al., 2010). Similar findings also have been found in AD (Schwindt and Black, 2009). Together, these studies support part of the atrophy-compensation hypothesis.

However, few studies have provided specific evidence for the association between smaller brain structures (e.g., gray matter density or volume loss) and higher brain activity in the PFC or LPC among aging adults. For example, Kalpouzos et al. (2012) found that the higher brain activity in PFC and LPC in older than younger adults during memory retrieval was eliminated after controlling for gray matter density in those regions, but not in other frontal and parietal regions (i.e., the left dorsomedial PFC and right LPC). This study provides both support for and against the general notion that atrophy should be associated with elevated brain activity. Similarly, Tyler et al. (2010) found that lower gray matter density in the left PFC and left temporal cortex was associated with higher activation in the right homologous regions during a language comprehension task in older adults. However, because gray matter density in each of those regions also was negatively correlated with brain activity in other frontal and temporal regions, these relationships appeared to be global rather than specific to nearby or contralateral regions. Other studies purportedly providing evidence for the atrophy-compensation hypothesis (1) assessed structure-function relationships only indirectly (Colcombe et al., 2005; Düzel et al., 2011; Persson et al., 2012; Thomsen et al., 2004), (2) found positive (not negative) structure-function relationships (Brassen et al., 2009; Rajah et al., 2011), (3) did not find a relationship between brain structure and function (Pudas et al., 2013), or (4) found relationships between brain function and white matter pathways using DTI (Daselaar et al., 2013; Persson et al., 2006), the latter of which can be difficult to classify as nearby or contralateral given the long distance of the fiber bundles. The lack of empirical support for the atrophy-compensation hypothesis is surprising given its widespread influence across multiple neurocognitive aging theories (Cabeza, 2002; Greenwood, 2007; Park and Reuter-Lorenz, 2009; Reuter-Lorenz and Cappell, 2008).

One possibility is that traditional measures of brain structure (e.g., gray matter density or volume loss) may not capture the type of atrophy that serves as a catalyst for neural compensation. FD, a measure of structural complexity, has been shown to be highly correlated with chronological age and a dementia diagnosis. FD quantifies fractal patterns, or irregularities, of cortical or subcortical surfaces similar to calculating the complexity of continental coastlines (Mandelbrot, 1967). One of the earliest studies using this method demonstrated that FD can provide better sensitivity to dementia-related differences in brain structure than thickness and gyrification (King et al., 2009, 2010). King et al. (2010) additionally found that FD was more strongly correlated with global cognition than other structural measures. In a series of studies, Madan and Kensinger extended this method to cognitively normal adults across the adult lifespan and also found higher correlations with chronological age and are more reliable in test-retest assessments than conventional measures (Madan and Kensinger, 2016, 2017b; see also Liu et al., 2020). Subsequent studies showed that these benefits can be found with improved precision in more localized regions, including cortical parcellations (Madan and Kensinger, 2018) and subcortical regions (Madan, 2019; Madan and Kensinger, 2017a).

The present study had 3 primary goals. The first goal was to provide a novel test for the basic premise of the atrophy-compensation hypothesis that lower brain integrity is associated with higher brain activity in a sample of middle-aged and older adults. We used FD as a proxy for brain integrity and task-related functional magnetic resonance imaging (fMRI) during encoding

and retrieval in a paired association task to assess potential negative associations with brain activity. In accordance with the atrophy-compensation hypothesis, lower FD should be associated with higher task-related brain activity in the PFC and LPC. We also predicted that any such negative associations would be stronger at retrieval than encoding because of the greater task demands during this phase (Mandzia et al., 2004; McDonough et al., 2013). All fMRI analyses attempted to minimize vascular confounds to the blood oxygen level dependent (BOLD) signal by using a participant-specific hemodynamic response function (Handwerker et al., 2004; Huettel et al., 2001) and scaling the contrasts using resting state fluctuation analyses (Kalcher et al., 2013; Kannurpatti et al., 2011). We conducted additional control analyses to ensure that potential negative structure-function relationships did not also occur in a sample of healthy younger adults—a sample in which atrophy does not yet occur. The second goal was to test whether individual differences in FD would be associated with higher risk for dementia using a cumulative dementia risk score in middle-aged and older adults (McDonough et al., 2019). Given the previous associations with FD and AD (King et al., 2009, 2010), we predicted that higher dementia risk would be associated with lower FD in the medial temporal lobe (MTL) and regions within the default mode network, consistent with previously found patterns of atrophy in AD (e.g., Buckner et al., 2005). The third goal was to link the structure-function patterns to cognition. Although finding a positive correlation between higher brain activity and better cognitive performance intuitively captures the notion of successful compensation (Cabeza et al., 2018), some perspectives suggest between-subject correlations are not necessary, especially in the cases of *attempted* rather than *successful* compensation (Dennis and Cabeza, 2012). To the extent that such relationships with cognition are indeed compensatory, then similar relationships should not also be present in a healthy young adult sample—a possibility that we tested.

2. Materials and methods

2.1. Participants

Sixty-seven participants aged 50–74 were drawn from the Alabama Brain Study on Risk for Dementia. Of these, 4 participants were not included in the FD analysis for the following reasons: (1) no functional data to correct for movement, (2) poor quality structural scans, and (3) cognitive data were unavailable. In addition, 3 more participants were not included in the task fMRI analysis because of residual movement artifact. Details from the study can be found in our earlier publication (McDonough et al., 2019). All participants were excluded if they had contraindicators for MRI, were left-handed, and had a prior diagnosis of any neurological condition, stroke, traumatic brain injury, claustrophobia, history of substance abuse. Potential participants were included if they were free of dementia as measured by the St. Louis University Mental Status (SLUMS; Tariq et al., 2006), spoke English fluently, were right-handed, and had at least one of the following self-reported risks for dementia: subjective memory complaints, less than a high school education, African American or Hispanic ethnographic category, mild head trauma, family history of AD, current diagnosis of hypertension or systolic blood pressure greater than 140 mmHg, current diagnosis or a family history of heart disease, current diagnosis of high total cholesterol, history or current use of smoking tobacco, current diagnosis or family history of diabetes, and body mass index greater than 30 kg/m² (Livingston et al., 2017; Perkins, 1997). An additional control group of 18 younger adults from age 20 to 30 was used to test key aspects of the atrophy-compensation hypothesis. This group had similar inclusion and exclusion criteria

with the exception that dementia risk and the SLUMS were not assessed. All participants gave informed consent using methods approved by the institutional review board at The University of Alabama. Vision was normal or corrected to normal using MR-compatible glasses or contact lenses. Demographic characteristics can be found in [Table 1](#).

2.2. Procedures

2.2.1. Overall study

Across 2 to 3 sessions, participants completed a cognitive and MRI battery. From the cognitive battery, we used the scaled word reading subtest from the Wide Range Achievement Test-4 to estimate premorbid intelligence quotient (IQ). The MRI session included scans in the following order: resting-state, memory encoding, T1-structural scan, resting-state, memory retrieval, and a visual-motor checkerboard task. The analyses here focus on the structural and task-related functional data.

2.2.2. Cognitive assessment

2.2.2.1. SLUMS examination. The SLUMS was used to characterize global cognition in the middle-aged and older adult sample ([Tariq et al., 2006](#)). The assessment consisted of 11 questions that measured a variety of cognitive domains including orientation, memory, attention, and executive functioning. The maximum number of points possible was 30. The SLUMS outperforms other common global measures of cognition such as the Mini-Mental Status Examination ([Folstein et al., 1975](#)).

2.2.2.2. Premorbid intelligence. The Wide Range Achievement Test-4 Word Reading subtest was administered to measure participants' reading skills via pronunciation ability of increasingly more difficult words. Pronunciation ability is commonly used to assess one's cognitive ability level before potential aging or pathologically related declines in ability ([Wechsler, 1944](#)). Each score was scaled using age norms.

2.2.3. fMRI memory task

The encoding phase consisted of 64 pairs of pictures of either a face-object or face-scene pairs for 3 seconds followed by a judgment of learning for 2.16 seconds. The intertrial interval ranged from 1.72 to 17.20 seconds. The encoding phase was divided into 2 runs lasting for about 8 minutes each. The memory test consisted of 64 trials lasting 5.16 seconds in a 5-alternative-choice test. A previously viewed face was presented with 2 previously viewed objects, 2 previously viewed scenes, and a “never seen” option. Of the 4 possible picture choices, 1 was the target and 3 were lures.

Table 1
Participant characteristics

	Older	Younger
	M (SD) or N (%)	M (SD) or N (%)
N ^a	63	18
Mean age	60.19 (6.78)	23.22 (3.04)
Sex (female)	41 (65%)	10 (56%)
Race (non-Hispanic white)	40 (63%)	11 (61%)
Years of education	14.00 (2.81)	14.89 (2.19)
Dementia risk score	4.44 (1.80)	–
SLUMS	26.46 (2.86)	–
Premorbid intelligence	101.30 (17.63)	107.61 (10.73)
fMRI memory accuracy % ^a	0.34 (0.12)	0.57 (0.08)

Key: SD, standard deviation; SLUMS, St. Louis University Mental Status.

^a fMRI data from 3 participants were unavailable because of in-scanner movement or artifact; Young adults were not asked for dementia risk information nor were given the SLUMS.

Because all options should have been familiar to participants, accurate responses relied on recollection processes. The “never seen” option specifically referred to not remembering the face, thus precluding participants from making a correct response without complete guessing. The intertrial intervals ranged from 1.72 to 10.32 seconds. The retrieval phase was divided into 2 runs lasting for about 5 minutes each.

2.2.4. fMRI checkerboard task

The purpose of this task was to obtain a measure of each participant's hemodynamic response function (HRF) in the occipital cortex that was time-locked to the onset of the visual stimulation. A reversing checkerboard was presented for 1 second in each of 20 repetitions over 2 minutes. When the checkerboard was presented, the participant was instructed to view the image and tap a button on the MR-compatible box that was provided to them. Intertrial intervals varied between 8 seconds and 16 seconds with an average of 12 seconds, during which the participant viewed a fixation cross.

2.2.5. Resting state scans

Two resting state scans were collected that each consisted of 175 volumes over a 5-minute span. Participants were told to close their eyes but not fall asleep.

2.3. MRI acquisition and preprocessing

A 3T Siemens PRISMA scanner at the UAB Civitan International Neuroimaging Laboratory was used to collect MRI scans. High-resolution T1-weighted structural MPRAGE scans were acquired using parallel acquisition acceleration type = GRAPPA; acceleration factor = 3, TR = 5000 ms, TE = 2.93 ms, TI 1 = 700 ms, TI 2 = 2030 ms, flip angle 1 = 4°, flip angle 2 = 5°, FOV = 256 mm, matrix = 240 × 256 mm², in-plane resolution = 1.0 × 1.0 mm². Structural images were processed using a surface-based processing stream provided by FreeSurfer v6.0 ([Fischl, 2012](#); <http://surfer.nmr.mgh.harvard.edu>), including bias-field correction, intensity normalization, skull-stripping ([Dale et al., 1999](#); [Segonne et al., 2004](#)). Following these steps, FreeSurfer segments of gray/white matter and pial surfaces were used to estimate distance between boundaries, which were checked for accuracy using Mindcontrol ([Keshavan et al., 2018](#)). Edits were made by extending white matter boundaries or removing non-brain tissue. Cortical regions of interest (ROIs) were defined using the Desikan-Killiany-Tourville (DKT) atlas ([Klein and Tourville, 2012](#)). Subcortical ROIs were defined using the FreeSurfer aseg atlas.

The aseg and DKT files from FreeSurfer were then submitted to the calcFD toolbox ([Madan and Kensinger, 2016, 2017a](#); <http://cmadan.github.io/calcFD>) in Matlab. This toolbox calculates the FD of 3-dimensional structures, including both cortical and subcortical structures. In fractal geometry, several approaches have been proposed to quantify the complexity of natural structures; the approach here calculates the Minkowski–Bouligand or Hausdorff dimension in 3-dimensional space ([Kennedy and Lin, 1986](#); [Mandelbrot, 1967](#)). This analysis has been validated and found to be reliable even after head motion ([Madan, 2018](#); [Madan and Kensinger, 2017b](#)). Briefly, FD is a scale-invariant measure of how complex a natural structure is and is particularly sensitive to shape information, rather than the ‘size’ of a structure, as is the case with volumetric measures—that is, volume, thickness, and surface area. In other words, 2 structures can be the same size, but differ in FD because of variations in the folding structure, texture (‘bumpiness’), or compactness (see [Madan, 2019](#)). Conversely, 2 structures could be different in size and still have the same FD. See [Madan and Kensinger \(2016\)](#) for visualizations of cortical surfaces that are relatively high and low in FD.

All functional scans used T2*-weighted EPI sequences (56 interleaved axial slices, 2.5 mm thickness, TR = 1720 ms, TE = 35.8 ms, flip angle = 73°, FOV = 260 mm, matrix = 104 × 104 mm, in-plane resolution = 2.5 × 2.5 mm², multiband acceleration factor = 4). The functional data were unwarped, coregistered to the structural scan, and spatially smoothed (8-mm FWHM kernel) using Statistical Parametric Mapping 12 (SPM12). The BOLD signal was then denoised using Multivariate Exploratory Linear Optimized Decomposition into Independent Components (Beckmann and Smith, 2004). The resulting spatiotemporal components were flagged using an in-house script that applied machine learning to frequency and temporal elements indicative of potential artifacts. The flagged components were then regressed from the BOLD signal, also using Multivariate Exploratory Linear Optimized Decomposition into Independent Components. The denoised data were then warped into a study template using Advanced Neuroimaging Tools (Avants and Gee, 2004). For the resting state data, additional preprocessing steps were used to prepare the data for the resting-state fluctuation amplitude (RSFA) analyses. AFNI's *proc.py* (Cox, 1996) was used to despike the time series, remove the first 3 TRs, band-pass filter the time series between 0.01 and 0.08 Hz, demean the time series, and to regress out motion parameters, their derivatives, and white matter signal from the time series.

2.4. Statistical analyses

2.4.1. FD analysis

Movement parameters from the functional scans were calculated using Artifact Detection Tools (ART; Mazaika et al., 2005), including mean framewise displacement and mean root squared realignment parameters. Although research using a large healthy sample of adults showed very little influence of head motion on FD estimations (Madan, 2018), a growing body of research has shown that in-scanner movement can have large effects on brain size estimates, in general (Alexander-Bloch et al., 2016; Savalia et al., 2017). Thus, we checked to see if this was the case in our data set. Preliminary correlations between FD and movement (framewise displacement and root mean squared estimates) ranged from −0.43 to 0.10 across ROIs, suggesting that more movement was related to smaller estimates of FD values. In light of this relationship, we regressed out these movement parameters from all ROIs before further analyses.

Next, we aimed to reduce the data set by conducting a factor analysis on the 76 ROIs (62 neocortical and 14 subcortical). To determine the optimal number of factors, we conducted a parallel analysis (Valle et al., 1999). A factor analysis was then conducted using oblimin rotation. Subsequent regression analyses were then conducted using these factor scores. Specifically, we investigated linear and quadratic effects of age, dementia risk, and their interaction separately for each FD factor. The dementia risk score consisted of a sum of 11 possible self-reported risk factors that could be identifiable by participants (see [Participants](#) section). A backward stepwise regression method was used to eliminate higher order interaction terms to preclude saturating the model using a *p*-value threshold of 0.10. In the second set of regression analyses, we investigated the extent that each factor, age, dementia risk, and their interactions predicted cognition. In these latter analyses, we included premorbid IQ as a covariate to control for lifelong cognition.

2.4.2. Task fMRI analysis

All fMRI analyses were analyzed under the assumptions of the general linear model (GLM). We first estimated each participant's HRF using a finite impulse response function to model task-evoked brain activity in response to the flashing checkerboard using SPM.

In this GLM, 12 basis functions were modeled across a 21-second window and outliers derived from ART were entered as nuisance regressors. At the first level, an F-test was calculated across the 12 time points at a liberal threshold of *p* = 0.05 using an inclusive mask of the occipital cortex. At the peak voxel within this mask, parameter estimates were extracted, which were used to represent each participant's HRF. These new HRFs were then entered into GLM analyses using SPM separately for encoding and retrieval using a stick function (setting duration to 0 seconds). For both memory encoding and retrieval, 2 trial types of interest were included as regressors: trials subsequently remembered (or correctly remembered) and trials subsequently forgotten (or inaccurately remembered). Other regressors of noninterest included 6 motion parameters, binary flags for outlying trials derived from ART, and 2 binary constants for mean session effects. A high-pass filter of 128 seconds was used. The encoding contrast (subsequently remembered > subsequently forgotten) and the retrieval contrast (correctly remembered > incorrectly remembered) were then scaled to account for cerebrovascular reactivity differences within the BOLD signal. This scaling process was carried out by dividing the contrast images by the mean RSFA image from the resting-state scans. RSFA analyses calculate the standard deviation of the BOLD signal. These analyses were implemented separately for each run and averaged together. Whole-brain contrasts can be found in [Supplemental Materials](#).

2.4.3. Structure-function group analysis

Because our group analyses were concerned with the potential effects of atrophy on brain activity, our analyses focused on middle-aged and older adults only. To test our primary hypothesis, regression analyses were conducted using both FD factors as predictors of the whole-brain beta maps in the same model, but separately for the encoding and retrieval contrasts. Regression analyses were conducted using FSL's randomise function (Winkler et al., 2014). Randomise uses a nonparametric approach and generates 5000 permutations to create a null distribution from which to make inferences. Data were demeaned before conducting the analyses. Threshold-free cluster enhancement was used to correct for the family-wise error rate at the 0.05 level for each regression analysis. This method of significance testing has been argued to be more sensitive than cluster-based and voxel-based thresholding (Smith and Nichols, 2009). In exploratory analyses, we also tested nonlinear effects by including 4 regressors in a new model (linear and quadratic effects of each FD factor). To strengthen the inferences from any null results found in the PFC (the site of our primary hypothesis), we conducted additional analyses using Bayesian correlation pairs under the hypothesis that a negative correlation would be found between FD and brain activity. Specifically, mean values for brain activity (correct > incorrect) were calculated for each of the 22 PFC ROIs from the DKT atlas and paired with either the FD value in the same hemisphere (i.e., left brain activity with left FD) or in the contralateral hemisphere (i.e., left brain activity with right FD). Bayes factor values closer to 0 indicate stronger evidence that the structure-function correlations are not negatively correlated. Finally, we conducted the same analyses in the young control group to ensure that the same relationships were not found.

To better understand how the FD factors and the associated functional clusters from the abovementioned regression related to cognition, partial least squares regression (PLS-R) was conducted using the ExPosition package in R (Beaton et al., 2014). We chose this method because it permits a single analysis that captures the shared variance across the inputted measures to explain most of the covariance in the data. For this analysis, 2 matrices were created: one that represented each brain measure (the FD factors and

clusters of brain activity) and one that represented cognition (global cognition via the SLUMS, memory accuracy, and premorbid IQ). The cross product of these 2 matrices was decomposed into mutually orthogonal latent variables using singular value decomposition. The latent variable scores represented the weights of the brain factors that contributed to higher or lower cognition for each test. Correlations between the latent variable scores from each X and Y matrix were used to determine significance. Additional analyses were carried out including age group (younger vs. middle-aged/older) in the model to test that these relationships did not also occur for younger adults.

3. Results

3.1. Factor analysis of FD across all ROIs

Among the middle-aged and older adults, a parallel factor analysis suggested 2 factors, which explained 29.0% and 13.0% of the variance for each factor. Factor loadings can be found in Fig. 1 and Supplemental Table 1. Similar loadings were found in the young adult control group that was included (Supplemental Table 2). For more details on the analyses, see Supplemental Materials. The first factor loaded on neocortical brain regions with the highest loadings consisting of lateral frontal and parietal regions. The second factor loaded subcortical brain regions with the highest loadings consisting of the pallidum, caudate, and putamen. Occipital and orbitofrontal cortices had the lowest loadings across the 2 factors, which are sometimes found to be spared with age using other structural measures (e.g., Fjell et al., 2009; Raz et al., 2005). No clear pattern of laterality was observed. The 2 factors were moderately correlated with one another ($r(61) = 0.37, p = 0.003$).

3.2. Associations between FD, age, and dementia risk

For factor 1 (neocortical structures), the age \times dementia risk interaction ($p = 0.31$) was not significant and so was removed from the analysis. In the next model, neither age ($p = 0.47$) nor dementia risk ($p = 0.36$) was significant. None of the quadratic effects of age, dementia risk, or their interaction were significant (p 's > 0.48). When the young adult control group was included, age ($b = -0.029, SE = 0.006, p < 0.001$) but not age² ($p = 0.75$) was significant. For factor 2 (subcortical structures), the age \times dementia risk interaction ($p = 0.46$) was not significant and so was removed from the analysis. In the next model, age was negatively related to FD

($b = -0.060, SE = 0.022, p = 0.0082$), but dementia risk was not ($p = 0.19$). None of the quadratic effects of age, dementia risk, or their interaction were significant (p 's > 0.54). Steiger's Z-test revealed that the strengths of these 2 linear age effects were marginally different from one another ($Z = 1.36, p = 0.091$). When the young adult control group was included, age ($b = -0.033, SE = 0.006, p < 0.001$) but not age² ($p = 0.49$) was significant. Together, the findings show that older age is associated with much lower cortical and subcortical FD than younger adults, but when focusing on middle-aged and older adults, differences are only found for subcortical structures accompanied by no effects of AD risk (see Fig. 2).

3.3. Associations between FD and brain activity during memory

To the extent that FD can be interpreted as a proxy for brain atrophy, lower FD should be associated with higher frontoparietal brain activity. In contrast to this prediction, neither factor 1 nor factor 2 was associated with frontoparietal brain regions during successful memory encoding. However, FD in the neocortex (factor 1) was negatively associated with brain activity during memory retrieval in several posterior brain regions, including bilateral precuneus, right supplementary motor area, right paracentral lobule, and left inferior parietal lobule (see Table 2 and Fig. 3). Of these regions, the largest cluster was in the precuneus, which is a major hub within the default mode network. No significant associations were found with FD in subcortical regions (factor 2) nor were associations found for quadratic associations between FD and brain activity. Correlations between brain activity and cortical FD in young adults revealed no significant relationships (all p 's > 0.08 , uncorrected) with 3 of the 4 clusters showing numerically positive relationships (ranging from $r = 0.04$ to 0.41). These nonsignificant relationships in the young control sample bolster the idea that the negative structure-function relationships are unique to middle-aged and older adults in which atrophy has started in some individuals.

Given the abovementioned null effects in the PFC among middle-aged and older adults, we conducted additional analyses to provide evidence for the null hypothesis using a Bayesian approach (see Fig. 4). For structure-function relationships in the same hemisphere ROIs, the Bayes factor scores ranged from 0.052 to 1.19. Across both encoding and retrieval, 29 of the 44 (66%) ROIs showed moderate to strong evidence for the null hypothesis, 13 (30%) showed anecdotal evidence for the null hypothesis, and 2 showed anecdotal evidence for a negative structure-function relationship. The 2 regions that showed anecdotal evidence in favor of a negative

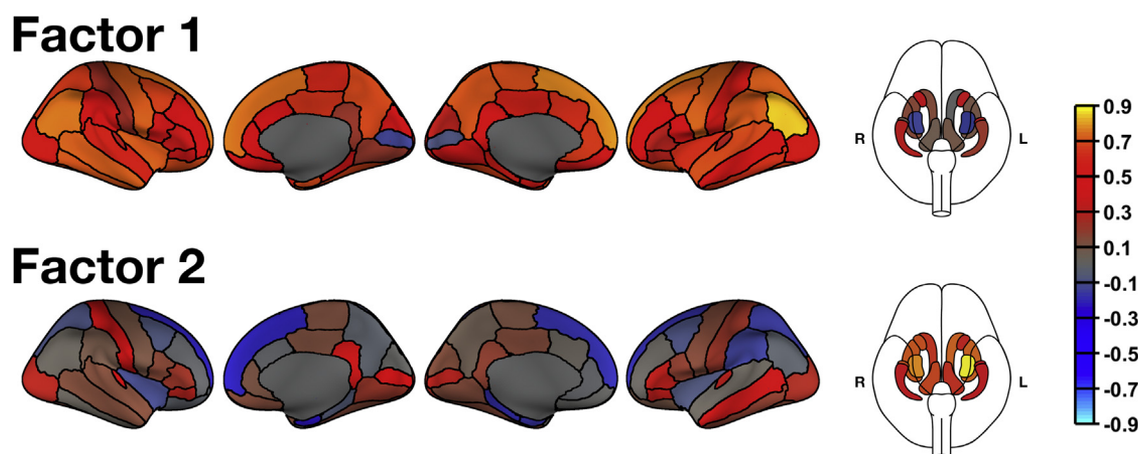


Fig. 1. Factor loadings plotted on the cortical surface and subcortical parcellations.

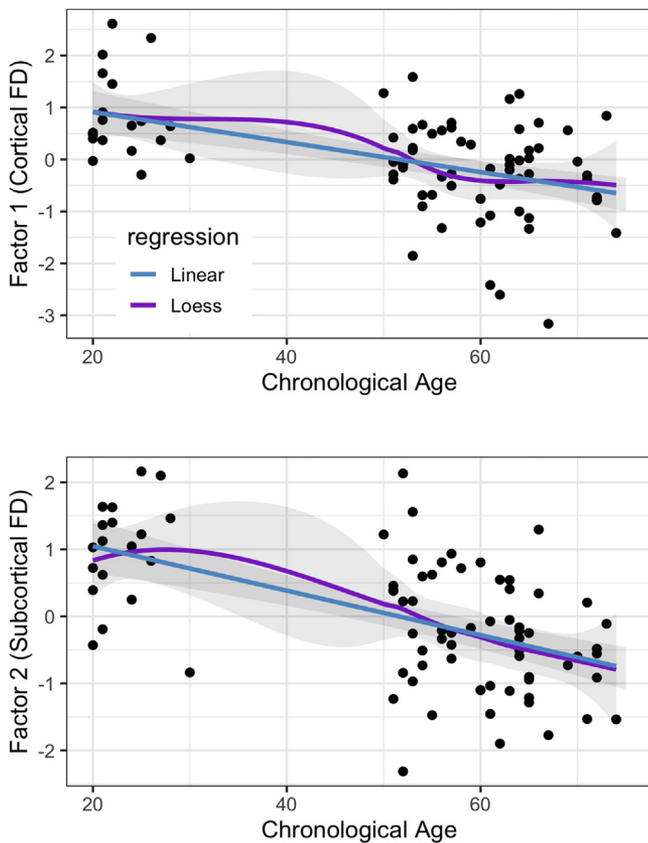


Fig. 2. Scatterplots illustrating the relationship between chronological age (x-axis) and fractal dimensionality (y-axis) for the first factor that represented fractal dimensionality in neocortex structures (upper panel) and the second factor that represented fractal dimensionality in subcortical structures (lower panel). In the middle-aged and older adult sample, only factor 2 revealed a significant relationship with age ($r = -0.29$, 95% CI $[-0.54, -0.05]$). However, when young adults were included in the models, both showed strong associations with age ($r = -0.49$, 95% CI $[-0.69, -0.30]$ for factor 1 and $r = -0.56$, 95% CI $[-0.74, -0.37]$ for factor 2). Blue lines represent a linear fit and the purple lines represent a non-linear fit with 95% confidence intervals in gray. Abbreviations: CI, confidence interval. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

structure-function relationship were the left pars triangularis during encoding (BF = 1.19) and the right pars opercularis during retrieval (BF = 1.07). For structure-function relationships in the contralateral hemisphere ROIs, the Bayes factor scores ranged from 0.064 to 0.92. Across both encoding and retrieval, 35 of the 44 (80%) ROIs showed moderate to strong evidence for the null hypothesis and 9 (20%) showed anecdotal evidence for the null hypothesis. These findings provide additional evidence against the atrophy-compensation hypothesis.

Table 2

MNI coordinates for association between fractal dimensionality and brain activity during memory retrieval (correct > incorrect)

MNI coordinates			Region	BA	T-value	Cluster size
x	y	z				
-28	-79	46	Left inferior parietal lobule	7	4.74	11
7	-46	69	Bilateral precuneus	7	4.44	83
8	-15	64	Right supplementary motor area	6	4.22	11
11	-24	73	Right paracentral lobule	6	4.01	55

Key: BA, Brodmann area; MNI, Montreal Neurological Institute. Clusters are labeled in order of statistical magnitude.

3.4. Associations between FD, brain activity, and cognition

Among middle-aged and older adults, the PLS PLS-R analysis revealed 2 significant components (see Fig. 5). The first latent variable explained 93.69% of the covariance among the variables, $r(58) = 0.31$, $p = 0.017$. This first latent variable revealed that lower FD and higher brain activity during retrieval were associated with worse cognition across all of the cognitive measures. The second latent variable explained 4.91% of the covariance among the variables, $r(58) = 0.36$, $p = 0.0049$. This latent variable revealed that individuals with better word reading (premorbid IQ) revealed lower brain activity in the left inferior parietal lobule during retrieval. Note that removing the FD factors from the analysis (thereby removing the redundancy from the previous structure-function analysis) results in the same 2 factors. For nonlinear effects on cognition, see [Supplemental Materials](#). These results provide evidence that higher FD and the related lower levels of brain activity are associated with better cognition, with stronger effects for neocortical FD than subcortical FD in middle-aged and older adults. In contrast, the young adult group revealed only one significant latent variable that explained 86.25% of the covariance among the variables, $r(16) = 0.60$, $p = 0.009$, such that higher cortical FD and higher brain activity associated with better cognition with the caveat that brain activity overall did not strongly contribute to the model at all (see [Supplemental Materials](#)). Given that the structure-function-cognition relationships were largely in the opposite direction than that found in the older sample, these relationships are likely not due to natural individual differences that occur throughout the lifespan but are unique to the older sample.

Given this negative relationship with brain activity and cognition in the first latent variable in the older sample, we tested whether Simpson's paradox might be at play ([Cabeza et al., 2018](#)). Assuming the atrophy-compensation hypothesis was correct, this paradox would predict that higher brain activity would be associated with higher cognition, but only for individuals with the greatest brain atrophy. In contrast, there might be no relationship (or a negative relationship) between brain activity and cognitive performance for those with the least atrophy. We tested this paradox by conducting a brain activity \times brain atrophy interaction on cognitive performance. Of the 3 cognitive measures, only the word reading (i.e., premorbid IQ) evidenced interactions in the right paracentral lobule (interaction $p = 0.011$) and the right supplementary motor area (interaction $p = 0.024$). However, the interaction was in the opposite direction than that predicted by Simpson's paradox (see Fig. 6). In individuals with the most brain atrophy (lower cortical FD), higher brain activity was associated with poorer word reading scores. In individuals with the lowest brain atrophy (higher cortical FD), higher brain activity was associated with higher word reading scores.

4. Discussion

The present study revealed 2 separable patterns of FD, a measure of structural complexity. One pattern was more strongly associated with cortical structures and the other more strongly associated with subcortical structures. Replicating previous research, lower FD factor values were associated with advanced age ([Madan, 2019](#); [Madan and Kensinger, 2017a](#)). However, we also found that among middle-aged and older adults, only subcortical FD was associated with age, indicating that cortical FD decreases earlier in the lifespan (i.e., middle age), whereas subcortical FD continues to decline into old age. In contrast to these age effects, no relationship between FD and dementia risk

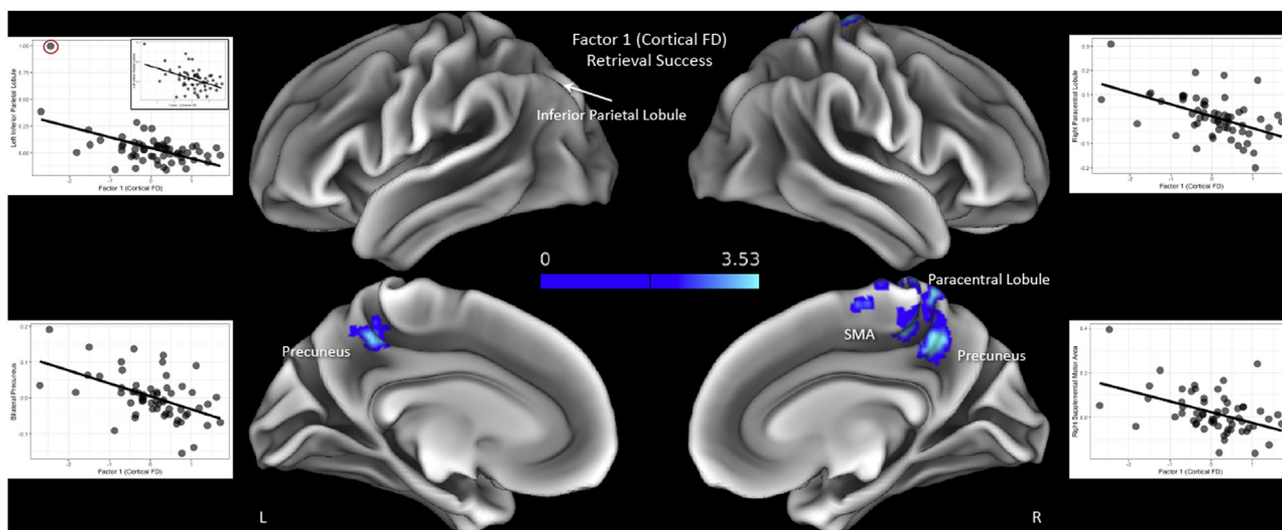


Fig. 3. Depiction of the negative association between fractal dimensionality in the neocortex (factor 1) and successful memory retrieval on the cortical surface. Scatterplots are shown in either side of the images with cortical FD on the x-axis and change in BOLD signal on the y-axis. The scatterplot inset in the upper left corner for the left inferior parietal lobule shows plot without the circled outlier ($r = -0.43$, p -value = 0.00060). Note that the left inferior parietal lobule was significant, but is not clearly shown in the surface rendering. Abbreviations: BOLD, blood oxygen level dependent; FD, fractal dimensionality; L, left; R, right.

was found. Beyond characterizing these basic relationships with FD, we critically provided a novel test for the atrophy-compensation hypothesis, which serves as a key principle found in some current neurocognitive theories of aging (Cabeza, 2002; Greenwood, 2007; Park and Reuter-Lorenz, 2009; Reuter-Lorenz and Cappell, 2008). This hypothesis proposes that brain atrophy causes a compensatory neural response in the form of increased brain activity in the PFC and LPC likely stemming from the recruitment of executive control functions during other cognitive tasks, including episodic memory. Although we did find support

for the general notion that lower brain structure is associated with more brain activity uniquely in our older sample, most of these associations were not found within the PFC and LPC. Instead, most of these negative associations were found in the medial parietal cortex, particularly in bilateral precuneus. Finally, we found that this atrophy-compensation pattern was most dominantly associated with poorer cognition in middle-aged and older adults. Young adults did not reveal a strong relationship between brain activity and cognition in the same brain regions. We elaborate on the major findings in more detail in the following sections.

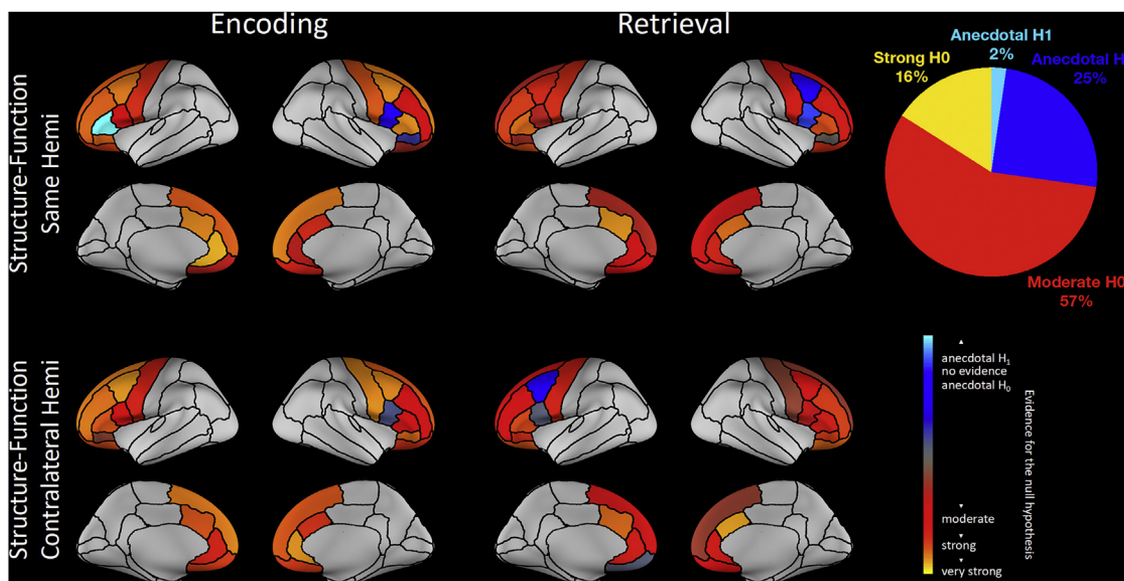


Fig. 4. Results from the Bayes correlation analysis between FD and brain activity showing a brain map of Bayes factor scores within prefrontal cortex regions of interest (ROIs) for memory encoding (left) and memory retrieval (right) in the same hemisphere (top) and the contralateral hemisphere (bottom). Bayes factor scores approaching 0 (from gray to red to yellow) indicate evidence for the null hypothesis (a negative structure-function relationship does not exist), whereas values greater than 1 (in blue) show anecdotal (i.e., weak) evidence for a negative association. Most of the ROIs showed moderate to strong evidence for the null hypothesis, whereas one-quarter of the ROIs showed anecdotal evidence for the null hypothesis and only 2% of the ROIs showed anecdotal evidence for a negative-structure function relationship. Abbreviations: FD, fractal dimensionality. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

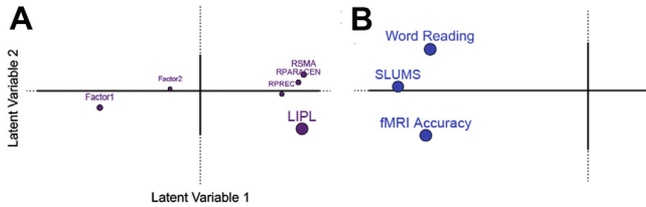


Fig. 5. Scatterplots showing the results of the partial least squares regression analyses assessing the association between brain measures and cognitive measures. (A) The left panel shows how the brain measures are expressed across the first latent variable (x-axis) and the second latent variable (y-axis). The axes represent the weights of each of the latent variable such that increasing weights move further away from the center of the axis (0, 0). The diameters of the circles are proportionate to the amount of total covariance explained by that factor. (B) The right panel shows how the cognitive measures are expressed across the first latent variable (x-axis) and the second latent variable (y-axis). Latent variable 1 shows that lower FD values and higher retrieval-related activity were associated with poorer cognition. Latent variable 2 shows that lower brain activity in the left parietal cortex was associated with better word reading ability (i.e., premorbid IQ) and lower memory accuracy. Abbreviations: FD, fractal dimensionality; LIPL, left inferior parietal lobule; RPARACEN, right paracentral lobule; RPREC, right precuneus; RSMA, right supplementary motor area; SLUMS, St. Louis University Mental Status.

4.1. Differential sensitivity of FD in the aging brain and cognition

In an attempt to simplify the FD analyses, a factor analysis was conducted to find groups of brain regions that covaried together. We found 2 very clear patterns that separated FD into cortical and subcortical factors. These differences might be interpreted in several ways. One methodological interpretation is that the calculation for FD on volumetric shapes systematically differs from the calculation for gyri, potentially due to the different biomechanical constraints on the relative growth/cohesion of tissue in the respective regions. Given the qualitatively different shapes of the 2, it should not be a surprise that analytical properties might differ as well.

Another interpretation is that the biological significance of these 2 sets of regions differs in meaningful ways, at least in the current sample. Consistent with this latter perspective, chronological age had a stronger association with FD in subcortical structures than cortical structures, but brain activity during memory retrieval and cognition had a stronger association with cortical structures than subcortical structures (as summarized in Fig. 4). Although not directly compared in past research, prior studies have generally found stronger correlations between age and subcortical FD than

cortical FD (Madan, 2019; Madan and Kensinger, 2016, 2017a). These findings suggest that FD in cortical regions should be considered separately from FD in subcortical regions when trying to understand age differences in brain morphology.

Only 2 other studies to date have related FD to cognition. The first did so in older adults with mild AD and healthy controls (King et al., 2010). When collapsing across both groups, lower global cognition as measured by the ADAS-Cog was associated with lower FD scores in the neocortex (i.e., a positive relationship)—although only global cortical FD was examined; no regional parcellation was considered. More recently, Liu et al. (2020) examined longitudinal changes in FD in older adults aged 70–90 year old. This study also assessed FD and global cognition. In their analyses, they found lobe-wise cortical parcellations of FD were positively associated with global cognition in the temporal lobe, occipital lobe, and several subcortical structures. They further found that FD partially mediated age-related changes in global cognition declines, with many of the effects stronger than the mediating effects of cortical thickness. Extending these previous results, the present study shows that even in young and middle-aged adults, higher cortical FD is associated with better global cognition and other cognitive domains including premorbid IQ and associative memory.

4.2. Testing the atrophy-compensation hypothesis

Madan and Kensinger (2016, 2018); Madan (2019) showed that FD is more correlated with chronological age than traditional measures of structural indices captured by T1 MRIs, including volume, cortical thickness, and gyrification. Similarly, FD can better distinguish between older adults with mild AD than healthy controls compared with cortical thickness and gyrification indices (King et al., 2010). More recent studies have demonstrated age-related decreases in FD longitudinally as well (Liu et al., 2020; Madan, 2020).

With this idea in mind, we tested the extent that using FD in conjunction with measures of brain activity could provide evidence for the hypothesis that lower brain integrity causes a compensatory neural response in the form of higher brain activity in the PFC and LPC (i.e., the atrophy-compensation hypothesis). We focused this analysis on middle-aged and older adults who would have the most likely of presenting with variable degrees of atrophy. Notably, the first FD factor most highly loaded onto the PFC and LPC, thus providing a strong test for this hypothesis. We found no evidence for such associations during memory encoding and found only small

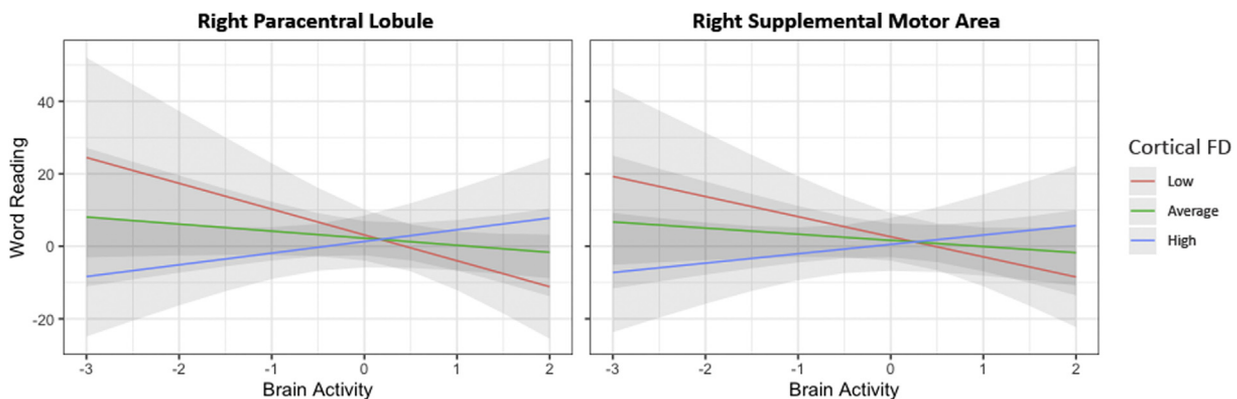


Fig. 6. Marginal means plots testing whether brain activity-cognition relationships differed between those with low atrophy (high fractal dimensionality) in blue, average atrophy in green, and high atrophy (low fractal dimensionality) in red. The left panel shows the significant brain structure \times brain activity interaction in the right paracentral lobule, and the right panel shows the significant interaction in the right supplementary motor area. For both regions, higher brain activity was associated with poorer word reading (i.e., premorbid IQ) in adults with higher atrophy, whereas higher brain activity was associated with better word reading in adults with lower atrophy. The gray areas represent 95% confidence intervals. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

evidence in support of this hypothesis during memory retrieval. Although several parietal regions evidenced this inverse relationship in our middle-aged and older adult sample (and not the young adult controls), only one small cluster in the LPC was negatively associated with the cortical FD factor. However, given the small size of the cluster and that most studies emphasize the role of the PFC, specifically, in such compensatory processes, this finding does not provide strong support for the atrophy-compensation hypothesis. Bayesian analyses directly targeting a negative structure-function relationship in the PFC provided moderate to strong evidence in nearly every ROI for an absence of such a relationship. Thus, the present findings add to an already mixed literature against a strict interpretation of the atrophy-compensation hypothesis (e.g., Brassens et al., 2009; Pudas et al., 2013; Rajah et al., 2011).

At least in the current paradigm, the present findings suggest that neural compensation may not always occur in the PFC and LPC. One possibility is that the PFC is less flexibly used in the presence of atrophy than expected. Intervention studies can provide causal evidence of the role that the PFC might play in compensating for cognitive abilities. McDonough et al. (2015) tested whether challenging activities over a 14-week period would increase the engagement of frontal-parietal brain regions involved in initiating and sustaining effortful cognitive processes. In contrast to predictions, only a small portion of the precentral gyrus (near the frontal eye fields) showed increased brain activity after the intervention, which did not differ from the active control groups. Instead, the mid-cingulate cortex, medial parietal cortex, and LPC showed reliable increases in activity after the intervention that differed from the control groups. The findings from this intervention are quite consistent with the present study and offer some preliminary evidence that the lateral PFC may not always be the primary source of neural compensation.

Another possibility is that neural compensation might occur anywhere near the site of the atrophy. For example, Daselaar et al. (2013) found that older adults with poorer white matter integrity in the PFC had higher brain activity in a nearby gray matter region, whereas those with poorer white matter integrity in the MTL had higher brain activity in a nearby gray matter region. The impact on cognition from this more generic pattern of “less wiring, more firing” was specific to the brain regions involved. They found that the stronger atrophy-compensation relationship in the PFC was associated with poorer executive function, whereas the stronger atrophy-compensation in the MTL was associated with poorer memory function. These authors argued that older adults successfully compensated for the brain atrophy because brain activity was defined as correct memory relative to incorrect memory and rejected the notion that successful compensation should be defined by interindividual differences (cf. Wang and Cabeza, 2016). A similar idea is that the spatial relationships between structure and function are only tightly coupled in unimodal sensory areas but are decoupled in transmodal regions like the PFC because of the hierarchical organization of the brain (Suarez et al., 2020). Both of these related perspectives suggest that atrophy in one region may be associated with brain function in different regions. Our findings are more aligned with this perspective because the cortical FD factor loaded highly on the parietal cortex, which was the site that a negative association was found. In addition, our results can be interpreted as successful compensation for 2 reasons. First, lower cortical FD was associated with higher correct than incorrect brain activity, thereby associating this pattern with successful memory performance. Second, this structure-function pattern was unique to middle-aged and older adults; young adults did not demonstrate these relationships, which would be expected if older, but not younger adults needed to engage in compensatory processes. Given that lateral and medial parietal cortices often are recruited during

memory retrieval (e.g., Rugg and Vilberg, 2013; Spaniol et al., 2009) in young adults, the type of compensation would be categorized as upregulation because the same regions used in younger adults are being upregulated in this sample (Cabeza et al., 2018).

Although our findings might be interpreted as successful compensation, not all researchers agree on how to characterize compensation. Many researchers believe that higher brain activity should be considered compensatory if higher activity were positively associated with individual differences in cognition (Cabeza et al., 2018). In the present study, we found that individuals who recruited more brain activity were *less* likely to remember correct associations, overall. From this interindividual differences perspective, the present findings provide behavioral evidence for brain dysfunction such as neural inefficiency (e.g., McDonough et al., 2013) or even neural toxicity (Pasquini et al., 2019).

These 2 perspectives might be reconciled by considering the effects of both the latent variables from the PLS-R analysis. Given that interindividual differences often are larger and can mask intraindividual differences (Aguirre et al., 1998; Leontiev and Buxton, 2007), one might consider the first latent variable as representing such large interindividual differences and the second latent variable as representing the residual intraindividual differences. Thus, the present findings might be interpreted such that middle-aged and older adults with higher lifelong cognition (estimated across the 3 cognitive measures) have generally (1) higher structural complexity and (2) more efficient (i.e., lower) brain activity. This interpretation is consistent with theories of brain maintenance (Cabeza et al., 2018; Nyberg et al., 2012), in which those adults who are able to maintain high structural and functional integrity also are able to maintain high levels of cognition. A related interpretation is that these individual differences are related to traits stable across the adult lifespan. However, the second latent variable dissociates premorbid IQ and fMRI memory accuracy. Thus, in the context of the specific memory retrieval task, higher structural complexity in the neocortex and higher fMRI activity in the left inferior parietal lobule are associated with better memory performance. This positive (rather than negative) association between the 3 modalities (structure, function, and cognition) would be predicted by attentional theories of memory (Cabeza et al., 2008; Ciaramelli et al., 2008). Note that, in the second latent variable, brain activity in the precuneus did not contribute to fMRI performance (as inferred by the centroid being close to the center of the y-axis). In fact, the part of the precuneus implicated in this study is more dorsal than that found in many prior memory studies (e.g., Gilmore et al., 2015; Huijbers et al., 2012; Spaniol et al., 2009) and instead might be associated with other cognitive processes (Power et al., 2011; Yeo et al., 2011). Regardless of the role that this part of the precuneus might play in memory retrieval, the PLS-R analysis might be able to offer a new perspective that accounts for how structure-function relationships can be both positively and negatively related to cognition. Although the middle-aged and older adult samples provide support for this new perspective, the control analyses in younger adults only partly support this view. In younger adults, higher cortical structural complexity was associated with better cognition, but these patterns were not associated with “more efficient” brain activity, thus questioning whether all the individual differences in the first latent variable were due to lifelong relationships between brain activity and cognition.

4.3. Strengths and limitations

One large difference between this study and the other studies investigating fMRI is that we calibrated the BOLD signal in each individual. Most older adults, and especially those with poorer health associated with dementia risks, have changes in brain

vasculature that can cause alterations of the BOLD signal and lead to both false positives and false negatives (for review see, [Wright and Wise, 2018](#)). We calibrated the BOLD signal in 2 practical ways: we used a participant-specific HRF and scaled the contrasts using RSFA analyses. At least some prior claims of higher frontally-based activity might be due to vascular alterations rather than neural differences, possibly explaining why we failed to find evidence for the atrophy-compensation hypothesis in the PFC. The present study also tested the atrophy-compensation hypothesis during both memory encoding and retrieval, which offered 2 different tests of the hypothesis, albeit only within an episodic memory paradigm. Finally, the atrophy-compensation hypothesis is most relevant to aging adults who might be experiencing atrophy and so any relationships found should not also be found in healthy young adults—who would not be experiencing atrophy and therefore would not need to recruit compensatory mechanisms. We demonstrated that the same structure-function-cognition relationships were not found in younger adults.

These strengths were accompanied by limitations. The most pressing limitation is the cross-sectional nature of the sample that precluded longitudinal measures of atrophy. Thus, our measure of structural complexity (i.e., FD) is best considered a combination of lifelong brain complexity mixed with longitudinal changes in complexity. As FD has only recently been used to study aging, it has only very recently been examined in longitudinal studies ([Liu et al., 2020](#); [Madan, 2020](#)), and even then, comparisons with other markers of atrophy need to be explored further. Relatedly, we cannot tease apart the causal direction of our structure-function relationships. For example, the scaffolding theory of cognitive aging (STAC; [Park and Reuter-Lorenz, 2009](#)) initially proposed that structural changes caused functional changes, but was later revised in STAC-R ([Reuter-Lorenz and Park, 2014](#)) to have bidirectional influences, opening the possibility that earlier changes in brain function might cause structural changes. From this perspective, subtle changes in synaptic functioning as measured by fMRI might be detected earlier than the larger synaptic alterations sensitive to T1 MRI scans ([Mondadori et al., 2006](#); [Mosconi et al., 2007](#)). Another limitation is that the sample did not contain adults aged 75 years or older, thus providing sensitivity only for early alterations in brain structure and function and cannot speak to “old-old” samples (e.g., [Song et al., 2016](#)).

We argue that the present failure to find even moderate negative structure-function relationships in the PFC is consistent with the weak existing evidence for the atrophy-compensation hypothesis using other measures of brain structure that we outlined in the Introduction. However, our test is limited by the specific episodic memory task and contrasts used. On the one hand, many of the theoretical perspectives that include an atrophy-compensation principle are agnostic to the actual task or cognitive domain used. These theories propose that older adults need to recruit executive function processes regardless of the original task (e.g., original processes + executive function). On the other hand, had a different task been used, the specific brain regions that showed significant relationships between FD and brain activity likely would have differed, thereby revealing other relationships. Having said that, using an episodic memory task is critical because older adults consistently show poorer associative episodic memory, and studies investigating age-related differences in episodic memory often invoke the notion of neural compensation, thus providing an important context in which to test these ideas.

5. Conclusions

The present study highlights an understudied analysis of brain structure, FD, that has promise to reveal new insights into

morphological brain differences in the aging process. Using FD, we tested a key principle in some cognitive neuroscience theories of aging: lower brain structure should be associated with higher brain activity in nearby or contralateral brain regions, especially in the PFC. We found the predicted negative associations that were unique to middle-aged and older adults, but not in the PFC. The null results in the PFC were supported by Bayes factor evidence values. Therefore, this study provides a growing body of evidence that contributes to the falsification of the standard atrophy-compensation hypothesis. The past decade has been accompanied by the generation of multiple neurocognitive theories of aging. Many have been quite influential in shaping the aging narrative in the field, but it has been difficult to provide falsifiable evidence for any of these theories. Indeed, the failure to find such relationships has been attributed to lacking insufficient variability or lacking statistical power ([Reuter-Lorenz and Cappell, 2008](#)). Even evidence showing the opposite than expected relationship has been attributed to Simpson's paradox, suggesting that the direction of the relationship at the population level may be different from the direction at the level of subgroup or individual (see ([Cabeza et al., 2018](#))). At least in the present study, we found strong evidence against Simpson's paradox. To the extent that future studies also fail to find atrophy-compensations associated with the PFC as in the present study, these cognitive aging theories would need to be refined.

Disclosure statement

The authors declare no competing financial interests.

CRediT authorship contribution statement

Ian M. McDonough: Conceptualization, Data curation, Formal analysis, Funding acquisition, Investigation, Methodology, Project administration, Resources, Supervision, Validation, Visualization, Writing - original draft, Writing - review & editing. **Christopher R. Madan:** Formal analysis, Methodology, Resources, Software, Visualization, Writing - original draft, Writing - review & editing.

Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at <https://doi.org/10.1016/j.neurobiolaging.2020.10.023>.

References

- Aguirre, G.K., Zarahn, E., D'esposito, M., 1998. The variability of human, BOLD hemodynamic responses. *Neuroimage* 8, 360–369.
- Alexander-Bloch, A., Clasen, L., Stockman, M., Ronan, L., Lalonde, F., Giedd, J., Raznahan, A., 2016. Subtle in-scanner motion biases automated measurement of brain anatomy from in vivo MRI. *Hum. Brain Mapp.* 37, 2385–2397.
- Avants, B., Gee, J.C., 2004. Geodesic estimation for large deformation anatomical shape averaging and interpolation. *Neuroimage* 23, S139–S150.
- Beaton, D., Fatt, C.R., Abdi, H., 2014. An ExPosition of multivariate analysis with the singular value decomposition in R. *Comput. Stat. Data Anal.* 72, 176–189.
- Beckmann, C.F., Smith, S.M., 2004. Probabilistic independent component analysis for functional magnetic resonance imaging. *IEEE Trans. Med. Imaging* 23, 137–152.
- Brassen, S., Büchel, C., Weber-Fahr, W., Lehmbeck, J.T., Sommer, T., Braus, D.F., 2009. Structure–function interactions of correct retrieval in healthy elderly women. *Neurobiol. Aging* 30, 1147–1156.
- Buckner, R.L., Snyder, A.Z., Shannon, B.J., LaRossa, G., Sachs, R., Fotenos, A.F., Sheline, Y.L., Klunk, W.E., Mathis, C.A., Morris, J.C., Mintun, M.A., 2005. Molecular, structural, and functional characterization of Alzheimer's disease: evidence for a relationship between default activity, amyloid, and memory. *J. Neurosci.* 25, 7709–7717.
- Cabeza, R., 2002. Hemispheric asymmetry reduction in older adults: the HAROLD model. *Psychol. Aging* 17, 85.
- Cabeza, R., Dennis, N.A., 2012. *Frontal Lobes and Aging. Principles of Frontal Lobe Function*, 2d ed. Oxford University Press, New York, pp. 628–652.

- Cabeza, R., Albert, M., Belleville, S., Craik, F.I., Duarte, A., Grady, C.L., Lindenberger, U., Nyberg, L., Park, D.C., Reuter-Lorenz, P.A., Rugg, M.D., Steffener, J., Rajah, M.N., 2018. Maintenance, reserve and compensation: the cognitive neuroscience of healthy ageing. *Nat. Rev. Neurosci.* 19, 701–710.
- Cabeza, R., Ciaramelli, E., Olson, I.R., Moscovitch, M., 2008. The parietal cortex and episodic memory: an attentional account. *Nat. Rev. Neurosci.* 9, 613.
- Ciaramelli, E., Grady, C.L., Moscovitch, M., 2008. Top-down and bottom-up attention to memory: a hypothesis (AtoM) on the role of the posterior parietal cortex in memory retrieval. *Neuropsychologia* 46, 1828–1851.
- Colcombe, S.J., Kramer, A.F., Erickson, K.L., Scalf, P., 2005. The implications of cortical recruitment and brain morphology for individual differences in inhibitory function in aging humans. *Psychol. Aging* 20, 363.
- Cox, R.W., 1996. AFNI: software for analysis and visualization of functional magnetic resonance neuroimages. *Comput. Biomed. Res.* 29, 162–173.
- Dale, A.M., Fischl, B., Sereno, M.I., 1999. Cortical surface-based analysis: I. Segmentation and surface reconstruction. *Neuroimage* 9, 179–194.
- Daselaar, S.M., Iyengar, V., Davis, S.W., Eklund, K., Hayes, S.M., Cabeza, R.E., 2013. Less wiring, more firing: low-performing older adults compensate for impaired white matter with greater neural activity. *Cereb. Cortex* 25, 983–990.
- Düzel, E., Schütze, H., Yonelinas, A.P., Heinze, H.J., 2011. Functional phenotyping of successful aging in long-term memory: preserved performance in the absence of neural compensation. *Hippocampus* 21, 803–814.
- Fischl, B., 2012. Freesurfer. *Neuroimage* 62, 774–781.
- Fjell, A.M., Westlye, L.T., Amlie, I., Espeseth, T., Reinvang, I., Raz, N., Agartz, I., Salat, D.H., Greve, D.N., Fischl, B., Dale, A.M., 2009. High consistency of regional cortical thinning in aging across multiple samples. *Cereb. Cortex* 19, 2001–2012.
- Fjell, A.M., McEvoy, L., Holland, D., Dale, A.M., Walhovd, K.B., Alzheimer's Disease Neuroimaging Initiative, 2014. What is normal in normal aging? Effects of aging, amyloid and Alzheimer's disease on the cerebral cortex and the hippocampus. *Prog. Neurobiol.* 117, 20–40.
- Folstein, M.F., Folstein, S.E., McHugh, P.R., 1975. "Mini-mental state": a practical method for grading the cognitive state of patients for the clinician. *J. Psychiatr. Res.* 12, 189–198.
- Gilmore, A.W., Nelson, S.M., McDermott, K.B., 2015. A parietal memory network revealed by multiple MRI methods. *Trends Cogn. Sci.* 19, 534–543.
- Greenwood, P.M., 2007. Functional plasticity in cognitive aging: review and hypothesis. *Neuropsychology* 21, 657–673.
- Handwerker, D.A., Ollinger, J.M., D'Esposito, M., 2004. Variation of BOLD hemodynamic responses across subjects and brain regions and their effects on statistical analyses. *Neuroimage* 21, 1639–1651.
- Huettel, S.A., Singerman, J.D., McCarthy, G., 2001. The effects of aging upon the hemodynamic response measured by functional MRI. *Neuroimage* 13, 161–175.
- Huijbers, W., Vannini, P., Sperling, R.A., Pennartz, C.M., Cabeza, R., Daselaar, S.M., 2012. Explaining the encoding/retrieval flip: memory-related deactivations and activations in the posteromedial cortex. *Neuropsychologia* 50, 3764–3774.
- Kalcher, K., Boubeta, R.N., Huf, V., Biswal, B.B., Baldinger, P., Sailer, U., Filzmoser, P., Kasper, S., Lamm, C., Lanzenberger, R., Moser, E., 2013. RESCALE: voxel-specific task-fMRI scaling using resting state fluctuation amplitude. *Neuroimage* 70, 80–88.
- Kalpourous, G., Persson, J., Nyberg, L., 2012. Local brain atrophy accounts for functional activity differences in normal aging. *Neurobiol. Aging* 33, 623–e1.
- Kannurpatti, S.S., Motes, M.A., Rypma, B., Biswal, B.B., 2011. Increasing measurement accuracy of age-related BOLD signal change: minimizing vascular contributions by resting-state-fluctuation-of-amplitude scaling. *Hum. Brain Mapp.* 32, 1125–1140.
- Kennedy, S.K., Lin, W.H., 1986. FRACT—a FORTRAN subroutine to calculate the variables necessary to determine the fractal dimension of closed forms. *Comput. Geosciences* 12, 705–712.
- Keshavan, A., Datta, E., McDonough, I.M., Madan, C.R., Jordan, K., Henry, R.G., 2018. Mindcontrol: a web application for brain segmentation quality control. *Neuroimage* 170, 365–372.
- King, R.D., George, A.T., Jeon, T., Hyman, L.S., Youn, T.S., Kennedy, D.N., Dickerson, B., Alzheimer's Disease Neuroimaging Initiative, 2009. Characterization of atrophic changes in the cerebral cortex using fractal dimensional analysis. *Brain Imaging Behav.* 3, 154–166.
- King, R.D., Brown, B., Hwang, M., Jeon, T., George, A.T., 2010. Alzheimer's Disease Neuroimaging Initiative. Fractal dimension analysis of the cortical ribbon in mild Alzheimer's disease. *Neuroimage* 53, 471–479.
- Klein, A., Tourville, J., 2012. 101 labeled brain images and a consistent human cortical labeling protocol. *Front. Neurosci.* 6, 171.
- Leontiev, O., Buxton, R.B., 2007. Reproducibility of BOLD, perfusion, and CMRO2 measurements with calibrated-BOLD fMRI. *Neuroimage* 35, 175–184.
- Li, S.C., Lindenberger, U., Sikström, S., 2001. Aging cognition: from neuromodulation to representation. *Trends Cogn. Sci.* 5, 479–486.
- Li, H.J., Hou, X.H., Liu, H.H., Yue, C.L., Lu, G.M., Zuo, X.N., 2015. Putting age-related task activation into large-scale brain networks: a meta-analysis of 114 fMRI studies on healthy aging. *Neurosci. Biobehav. Rev.* 57, 156–174.
- Liu, H., Liu, T., Jiang, J., Cheng, J., Liu, Y., Li, D., Dong, C., Niu, H., Li, S., Zhang, J., Brodaty, H., 2020. Differential longitudinal changes in structural complexity and volumetric measures in community-dwelling older individuals. *Neurobiol. Aging* 91, 26–35.
- Livingston, G., Sommerlad, A., Orgeta, V., Costafreda, S.G., Huntley, J., Ames, D., Ballard, C., Banerjee, S., Burns, A., Cohen-Mansfield, J., Cooper, C., 2017. Dementia prevention, intervention, and care. *Lancet* 390, 2673–2734.
- Logan, J.M., Sanders, A.L., Snyder, A.Z., Morris, J.C., Buckner, R.L., 2002. Under-recruitment and nonselective recruitment: dissociable neural mechanisms associated with aging. *Neuron* 33, 827–840.
- Madan, C.R., 2018. Age differences in head motion and estimates of cortical morphology. *PeerJ* 6, e5176.
- Madan, C.R., 2019. Shape-related characteristics of age-related differences in subcortical structures. *Aging Ment. Health* 23, 800–810.
- Madan, C.R., 2020. Age-related decrements in cortical gyrification: evidence from an accelerated longitudinal dataset. *Eur. J. Neurosci.* <https://doi.org/10.1111/ejn.15039>.
- Madan, C.R., Kensinger, E.A., 2016. Cortical complexity as a measure of age-related brain atrophy. *Neuroimage* 134, 617–629.
- Madan, C.R., Kensinger, E.A., 2017a. Age-related differences in the structural complexity of subcortical and ventricular structures. *Neurobiol. Aging* 50, 87–95.
- Madan, C.R., Kensinger, E.A., 2017b. Test–retest reliability of brain morphology estimates. *Brain Inform.* 4, 107–121.
- Madan, C.R., Kensinger, E.A., 2018. Predicting age from cortical structure across the lifespan. *Eur. J. Neurosci.* 47, 399–416.
- Mandelbrot, B., 1967. How long is the coast of Britain? Statistical self-similarity and fractional dimension. *Science* 156, 636–638.
- Mandzia, J.L., Black, S.E., McAndrews, M.P., Grady, C., Graham, S., 2004. fMRI differences in encoding and retrieval of pictures due to encoding strategy in the elderly. *Hum. Brain Mapp.* 21, 1–4.
- Mazaika, P.K., Whitfield, S., Cooper, J.C., 2005. Detection and repair of transient artifacts in fMRI data. *Neuroimage* 26 (Suppl 1), S36.
- McDonough, I.M., Letang, S.K., Stinson, E.A., 2019. Dementia risk elevates brain activity during memory retrieval: a functional MRI analysis of middle aged and older adults. *J. Alzheimer's Dis.* 70, 1005–1023.
- McDonough, I.M., Wong, J.T., Gallo, D.A., 2013. Age-related differences in prefrontal cortex activity during retrieval monitoring: testing the compensation and dysfunction accounts. *Cereb. Cortex* 23, 1049–1060.
- McDonough, I.M., Haber, S., Bischof, G.N., Park, D.C., 2015. The synapse project: engagement in mentally challenging activities enhances neural efficiency. *Restor. Neurol. Neurosci.* 33, 865–882.
- Mondadori, C.R., Buchmann, A., Mustovic, H., Schmidt, C.F., Boesiger, P., Nitsch, R.M., Hock, C., Streffer, J., Henke, K., 2006. Enhanced brain activity may precede the diagnosis of Alzheimer's disease by 30 years. *Brain* 129, 2908–2922.
- Mosconi, L., Brys, M., Glodzik-Sobanska, L., De Santi, S., Rusinek, H., De Leon, M.J., 2007. Early detection of Alzheimer's disease using neuroimaging. *Exp. Gerontol.* 42, 129–138.
- Nyberg, L., Lövdén, M., Riklund, K., Lindenberger, U., Bäckman, L., 2012. Memory aging and brain maintenance. *Trends Cogn. Sci.* 16, 292–305.
- Park, D.C., McDonough, I.M., 2013. The dynamic aging mind: Revelations from functional neuroimaging research. *Perspect. Psychol. Sci.* 8, 62–67.
- Park, D.C., Reuter-Lorenz, P., 2009. The adaptive brain: aging and neurocognitive scaffolding. *Annu. Rev. Psychol.* 60, 173–196.
- Pasquini, L., Rahmani, F., Maleki-Balajoo, S., La Joie, R., Zarei, M., Sorg, C., Drzezga, A., Tahmasian, M., 2019. Medial temporal lobe Disconnection and hyperexcitability across Alzheimer's disease stages. *J. Alzheimers Dis. Rep.* 3, 103–112.
- Perkins, P., Annegers, J.F., Doody, R.S., Cooke, N., Aday, L., Vernon, S.W., 1997. Incidence and prevalence of dementia in a multiethnic cohort of municipal retirees. *Neurology* 49, 44–50.
- Persson, J., Nyberg, L., Lind, J., Larsson, A., Nilsson, L.G., Ingvar, M., Buckner, R.L., 2006. Structure–function correlates of cognitive decline in aging. *Cereb. Cortex* 16, 907–915.
- Power, J.D., Petersen, S.E., 2013. Control-related systems in the human brain. *Curr. Opin. Neurobiol.* 23, 223–228.
- Persson, J., Pudas, S., Lind, J., Kauppi, K., Nilsson, L.G., Nyberg, L., 2012. Longitudinal structure–function correlates in elderly reveal MTL dysfunction with cognitive decline. *Cereb. Cortex* 22, 2297–2304.
- Power, J.D., Cohen, A.L., Nelson, S.M., Wig, G.S., Barnes, K.A., Church, J.A., A.C., Laumann, T.O., Miezin, F.M., Schlaggar, B.L., Petersen, S.E., 2011. Functional network organization of the human brain. *Neuron* 72, 665–678.
- Pudas, S., Persson, J., Josefsson, M., de Luna, X., Nilsson, L.G., Nyberg, L., 2013. Brain characteristics of individuals resisting age-related cognitive decline over two decades. *J. Neurosci.* 33, 8668–8677.
- Rajah, M.N., Languay, R., Grady, C.L., 2011. Age-related changes in right middle frontal gyrus volume correlate with altered episodic retrieval activity. *J. Neurosci.* 31, 17941–17954.
- Raz, N., Lindenberger, U., Rodrigue, K.M., Kennedy, K.M., Head, D., Williamson, A., Gerstorff, D., Acker, J.D., 2005. Regional brain changes in aging healthy adults: general trends, individual differences and modifiers. *Cereb. Cortex* 15, 1676–1689.
- Reuter-Lorenz, P.A., Cappell, K.A., 2008. Neurocognitive aging and the compensation hypothesis. *Curr. Dir. Psychol. Sci.* 17, 177–182.
- Reuter-Lorenz, P.A., Park, D.C., 2014. How does it STAC up? Revisiting the scaffolding theory of aging and cognition. *Neuropsychol. Rev.* 24, 355–370.
- Rowe, J.W., Kahn, R.L., 1987. Human aging: usual and successful. *Science* 237, 143–149.
- Rugg, M.D., Vilberg, K.L., 2013. Brain networks underlying episodic memory retrieval. *Curr. Opin. Neurobiol.* 23, 255–260.
- Savalia, N.K., Agres, P.F., Chan, M.Y., Feczko, E.J., Kennedy, K.M., Wig, G.S., 2017. Motion-related artifacts in structural brain images revealed with independent estimates of in-scanner head motion. *Hum. Brain Mapp.* 38, 472–492.

- Schwandt, G.C., Black, S.E., 2009. Functional imaging studies of episodic memory in Alzheimer's disease: a quantitative meta-analysis. *Neuroimage* 45, 181–190.
- Ségonne, F., Dale, A.M., Busa, E., Glessner, M., Salat, D., Hahn, H.K., Fischl, B., 2004. A hybrid approach to the skull stripping problem in MRI. *Neuroimage* 22, 1060–1075.
- Smith, S.M., Nichols, T.E., 2009. Threshold-free cluster enhancement: addressing problems of smoothing, threshold dependence and localisation in cluster inference. *Neuroimage* 44, 83–98.
- Song, Z., McDonough, I.M., Liu, P., Lu, H., Park, D.C., 2016. Cortical amyloid burden and age moderate hippocampal activity in cognitively-normal adults. *Neuroimage* 12, 78–84.
- Spaniol, J., Davidson, P.S., Kim, A.S., Han, H., Moscovitch, M., Grady, C.L., 2009. Event-related fMRI studies of episodic encoding and retrieval: meta-analyses using activation likelihood estimation. *Neuropsychologia* 47, 1765–1779.
- Spreng, R.N., Wojtowicz, M., Grady, C.L., 2010. Reliable differences in brain activity between young and old adults: a quantitative meta-analysis across multiple cognitive domains. *Neurosci. Biobehav. Rev.* 34, 1178–1194.
- Suárez, L.E., Markello, R.D., Betzel, R.F., Misisic, B., 2020. Linking structure and function in macroscale brain networks. *Trends Cogn. Sci.* 24, 302–315.
- Tariq, S.H., Tumosa, N., Chibnall, J.T., Perry III, M.H., Morley, J.E., 2006. Comparison of the Saint Louis University mental status examination and the mini-mental state examination for detecting dementia and mild neurocognitive disorder—a pilot study. *Am. J. Geriatr. Psychiatry.* 14, 900–910.
- Thomsen, T., Specht, K., Hammar, Å., Nytingnes, J., Ersland, L., Hugdahl, K., 2004. Brain localization of attentional control in different age groups by combining functional and structural MRI. *Neuroimage* 22, 912–919.
- Tyler, L.K., Shafto, M.A., Randall, B., Wright, P., Marslen-Wilson, W.D., Stamatakis, E.A., 2010. Preserving syntactic processing across the adult life span: the modulation of the frontotemporal language system in the context of age-related atrophy. *Cereb. Cortex.* 20, 352–364.
- Valle, S., Li, W., Qin, S.J., 1999. Selection of the number of principal components: the variance of the reconstruction error criterion with a comparison to other methods. *Ind. Eng. Chem. Res.* 38, 4389–4401.
- Vincent, J.L., Kahn, I., Snyder, A.Z., Raichle, M.E., Buckner, R.L., 2008. Evidence for a frontoparietal control system revealed by intrinsic functional connectivity. *J. Neurophysiol.* 100, 3328–3342.
- Wang, W.C., Cabeza, R., 2016. Episodic memory encoding and retrieval in the aging brain. In: Cabeza, R., Nyberg, L., Park, D.C. (Eds.), *Cognitive Neuroscience of Aging: Linking Cognitive and Cerebral Aging*, fourth ed. Oxford University Press, pp. 301–336.
- Wechsler, D., 1944. *The Measurement of Adult Intelligence*. Williams and Wilkins, Baltimore.
- Winkler, A.M., Ridgway, G.R., Webster, M.A., Smith, S.M., Nichols, T.E., 2014. Permutation inference for the general linear model. *Neuroimage* 92, 381–397.
- Wright, M.E., Wise, R., 2018. Can blood oxygenation level dependent functional magnetic resonance imaging be used accurately to compare older and younger populations? A mini literature review. *Front Aging Neurosci.* 10, 371.
- Yeo, B.T.T., Krienen, F.M., Sepulcre, J., Sabuncu, M.R., Lashkari, D., Hollinshead, M., Roffman, J.L., Smoller, J.W., Zöllei, L., Polimeni, J.R., Fischl, B., 2011. The organization of the human cerebral cortex estimated by intrinsic functional connectivity. *J. Neurophysiol.* 106, 1125–1165.