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## Can Structural MRI Indices of Cerebral Integrity Track Cognitive Trends in Executive Control Function During Normal Maturation and Adulthood?

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## Abstract

We explored the relationship between structural neuroimaging-based indices of cerebral integrity and executive control function (ECF) in two groups of healthy subjects: A maturing group (33 subjects; 19–29 years) and a senescing group (38 adults; 30–90 years). ECF was assessed using the Executive Interview (EXIT) battery. Cortical indices of cerebral integrity included GM thickness, intergyral span, and sulcal span, each measured for five cortical regions per hemisphere. Subcortical indices included fractional anisotropy (FA), measured using track-based-spatial-statistics (TBSS), and the volume of T2-hyperintense WM (HWM). In the maturing group, no significant relationships between neuroanatomical changes and ECF were found; however, there were hints that late-term maturation of cerebral WM influenced variability in ECF. In the senescing group, the decline in ECF corresponded to atrophic changes in cerebral WM (sulcal and intergyral span) primarily in the superior frontal and anterior cingulate regions. A large fraction of the variability in ECF (62%) can be explained by variability in the structural indices from these two regions.

## Keywords

aging; morphology; cerebral health; atrophy; maturation

## INTRODUCTION

The upcoming decades will be marked by substantial increases in the proportion of population above 65 years of age [He et al., 2006]. This rapidly growing segment of the population is characterized by rising prevalence of cognitive impairment and disabilities. There is an urgent need for sensitive measures of the aging process to better understand the neurobiology of cognitive decline in this age group. Such measures may ultimately be used (1) for gauging factors that affect aging and neurodegenerative processes, (2) for assessment of genetic factors that predispose one to abnormal aging, and (3) for clinical trials that determine the therapeutic value of anti-dementia drugs and their ability to resist degenerative changes. A better understanding of cerebral aging requires a broad perspective that can be achieved by combining

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cognitive (behavioral) and noninvasive brain imaging methods, thereby providing data to guide development of comprehensive, neurobiologically realistic models of aging.

As an initial step in the development of such neurobiological models, we investigated whether neuroimaging-based indices of structural integrity could explain a significant portion of individual variation in executive control function (ECF). ECF was chosen because of its importance to normal cognitive functioning and dementia, as it facilitates planning and adaptation to novel situations, and regulates complex goal-driven actions [Collette et al., 2005, 2006]. Another advantage of using ECF is that associated neuroanatomical networks are generally well mapped. Lesion studies, and more recently functional imaging studies, have localized ECF and its deficits predominantly to frontal and cingulate regions [Collette et al., 2005, 2006; Royall et al., 2001].

Association between neuropsychological status and degree of cerebral atrophy has been studied in clinical populations affected with Alzheimer's disease, ischemic brain disorder, and multiple sclerosis [Csernansky et al., 2005; Du et al., 2005b; Ezekiel et al., 2004; Royall et al., 2001]. However, these disorders are generally characterized by more dramatic changes in neuropsychological status and in measurements of cerebral integrity. Finding an association between changes in neuropsychological measures with changes in regional cerebral integrity has been challenging in normal subjects because of the subtleness of these changes. Another challenge in studies of ECF in normal subjects is that ECF-sensitive measures follow a nonlinear, "U-shaped" trajectory with age. During cerebral maturation, the ECF improves with age reaching maturity levels in the 3rd–4th decades of life and then declines in later decades [Royall, 2000].

We used the following neuroimaging-based structural indices to study variability in ECF in healthy maturing and senescing individuals: Gray matter (GM) thickness, the width of intergyral span, the width of intrasulcal CSF spaces, fractional anisotropy (FA) of cerebral WM, and the volume of the T2-hyperintense white mater (HWM). Each anatomical measure are described below.

GM thickness is a neuroimaging marker of cortical integrity related to regional neuronal density in the cortical mantle [Jelsing et al., 2005; Lerch and Evans, 2005; Selemon et al., 1995]. Agerelated changes in cortical thickness are well described [Sowell et al., 2003]. In normal subjects, the average thickness of the gray matter mantle follows an inverted-U trajectory with age: GM thickness rises during maturation, reaching maximum during the 2nd and 3rd decades of life and then decreases linearly with age [Kochunov et al., 2007, 2008; Magnotta et al., 1999; Raz N et al., 1997]. The rate of cortical GM thickness reduction in abnormal aging could be different from that in healthy aging, and this difference could be an important diagnostic factor [Thompson et al., 2003, 2004].

Intergyral span is sensitive to changes in the volume of gyral WM that form the bulk of cortical gyri [Kochunov et al., 2005b]. Sulcal span is a measurement of dilatation of cortical sulci and the compensatory filling of the freed space by CSF. Age-related "widening" of cortical sulci was shown to be due to reduction in GM thickness and gyral WM volumes [Jernigan et al., 2001; Kochunov et al., 2005b; Magnotta et al., 1999; Symonds et al., 1999]. Visual assessments of regional "widening" of cortical sulci, specifically in the hippocampal, enthorhinal, and medial temporal cortices, are correlated with cognitive decline in mild cognitive impairment and dementia [Bastos Leite et al., 2004].

FA in cerebral WM is a sensitive index of regional breakdown of the microstructure of WM tracts. Global and regional decline in FA has been shown to be correlated with the progression and severity of various degenerative WM disorders such as multiple sclerosis, leukoaraiosis, and various dementias [Horsfield and Jones, 2002]. Previously we reported that FA values

decrease during normal aging and that whole-brain average FA values were highly correlated with other indices such as GM thickness and sulcal span [Kochunov et al., 2007]. We also observed that FA values of the thinly myelinated associative tracks had the greatest decline with age and the highest correlation with GM thickness and sulcal span [Kochunov et al., 2007].

HWM lesions, observed as regions of high signal intensity on T2-weighted MR studies, are often present in both normal aging individuals and patients affected with neurodegenerative disorders. HWM lesions arise from regions of accumulation of extra-cellular water associated with degradation of the myelin sheath of nonspecific etiology [Pantoni and Garcia, 1995]. The prevalence of HWM lesions has been reported to be as high as 60–100% in normal subjects 60 years old and older [de Leeuw et al., 2001, 2005]. These lesions appear at a greater extent in frontal regions [Fazekas et al., 2005; Raz et al., 2003] and their volume is correlated with global decreases in cerebral blood flow [Kraut et al., 2008; ten Dam et al., 2007], reduction in cerebral WM and GM volumes [Du et al., 2005a; Wen et al., 2006] and reduction in global and regional FA values [Kochunov et al., 2007].

In this study, we predicted that the five neuroimaging-based markers of structural integrity would partly explain intersubject variability in a sensitive executive control measure, the Executive Interview (EXIT) [Royall et al., 1992, 1994; Stokholm et al., 2005]. The EXIT provides a standardized assessment of ECF. It takes ~15 min to administer, has high internal consistency (r = 0.85), high inter-rater reliability (r = 0.90) [Chan et al., 2006; Royall et al., 2007] and is available in multiple language translations [Chan et al., 2006; Stokholm et al., 2005]. EXIT scores have been shown to correlate well with performance on other ECFsensitive tests, such as the Wisconsin Card Sorting Task (WCST) (r = 0.54), trail making part B (r = 0.64), Lezak's tinker toy test (r = 0.57), and the test of sustained attention (time, r = 0.64) 0.82; errors, r = 0.83). However, EXIT scores appear most strongly associated with cognitive disability and clinical outcomes [Lewis and Miller, 2007; Royall et al., 2003, 2007]. In normal aging populations, performance on EXIT was associated with both cross-sectional level of care [Royall et al., 2000, 2003], and longitudinal change in instrumental activities of daily living [Royall et al., 2004, 2005a]. Performance on EXIT was also shown to predict other clinically relevant outcomes such as medication compliance and medical decision making capacity [Allen et al., 2003; Dymek et al., 2001]. Important advantages of the EXIT include its brevity and its validity in older populations. Moreover, unlike the WCST, the EXIT has no appreciable learning effect, and is sensitive to change in repeated measures designs [Royall et al., 2005b]. This makes EXIT feasible for assessing and monitoring ECF in combination with neuroimaging-based measurements of cerebral integrity during studies of normal and neurodegenerative aging, treatment effects, and clinical interventions.

## **METHODS**

#### Subjects

Seventy one (44 females, 27 males), healthy, right handed, 19–90 years of age (average age =  $47.4 \pm 22.1$  years) were recruited as the part of the International Consortium on Brain Mapping (ICBM) project [Mazziotta et al., 1995,2001]. Subjects were treated as two groups with approximately equal numbers per group, maturing, younger (age 19–29, N = 33, 21/12 females/males), and senescing, older (age 30–90, N = 38, 23/12 females/males). Subject's medical history was reviewed to rule out endocrinal, neurological, and psychiatric illnesses. The study was performed with institutional review board approval, and subjects provided written informed consent. All neuropsychological testing and brain imaging was performed at the Research Imaging Center, University of Texas Health Science Center at San Antonio.

## MR Imaging

MR imaging was performed at the Research Imaging Center, UTHSC, using a Siemens 3T Trio scanner and a high-resolution 8-channel head coil.

**T1-weighted imaging**—High-resolution (isotropic 800 lm), T1-weighted images were acquired using a retrospective motion-corrected protocol [Kochunov et al., 2006]. Under this protocol, seven full-resolution volumes are acquired using a 3D Turbo-FLASH sequence with an adiabatic inversion magnetization preparation contrast pulse (scan parameters: TE/TR/ TI 5 3.04/2100/785 ms, flip angle = 11 degrees). The sequence yields uniform tissue contrast across the imaging volume. Imaging time was 26 min.

**Diffusion weighted imaging**—Single-shot, spin-echo, echo-planar gradient recalled echo, T2-weighted, sequence was used to acquire diffusion weighted data with the spatial resolution of  $1.7 \times 1.7 \times 3.0$  mm<sup>3</sup>. The sequence parameters were TE/TR = 87/ 8000ms, with 86 diffusion weighted directions and two diffusion weighing values of 0 and 700 s/mm<sup>2</sup>. The total sequence acquisition time was 12 min.

**T2-weighted imaging**—The T2-weighted data were acquired with a high-resolution (isotropic 1 mm), 3D turbo-inversion recovery Fluid Attenuated Inversion Recovery (FLAIR) sequence with the following parameters: TR/TE/TI/Flip angle = 5 s, 353 ms, 1.8 s, 180 degrees. This sequence uses a nonselective inversion recovery pulse to prevent CSF pulsation artifacts [Bakshi et al., 2000]. Imaging time was 15 min.

## Structural Image Processing

Structural image processing consisted of preprocessing of T1-weighted images, sulcal surface analysis, and processing of FA and HWM images.

**Preprocessing of T1-weighted images**—Preprocessing of T1-weighted images consisted of removing nonbrain tissues, global spatial normalization, and radio frequency (RF) inhomogeneity correction. Nonbrain tissues such as skin, muscle, and fat were removed using an automated skull stripping procedure provided by the brain extraction tool [Smith, 2002] (see Fig. 1A). The extracted brain images were cropped at the level of the cerebellum. Next all brain images were globally spatially normalized to the Talairach coordinate system to remove gross differences in brain location, size, and orientation. Automated, nine-parameter (three rotations, three scales, three translations) FSL FLIRT global spatial normalization software [Smith et al., 2004] was used to register all images to a Talairach template [Kochunov et al., 2002]. All images were resliced to isotropic 800 lm spacing using a 3D, 15-voxel-wide sinc interpolation kernel. Images were radio frequency (RF) inhomogeneity corrected using the FAST tool (Fig. 1B) [Zhang et al., 2001]. Default FAST parameters were used. RF corrected images were imported into the BrainVISA database and processed through its sulcal extraction and identification pipelines, as described by Kochunov et al., [2005].

**Sulcal surface processing**—The BrainVISA image processing pipeline extracts hemispheric spherical meshes from segmented gray and white matter images (Fig. 1C,D). The medial surface of the cortical in-folds is obtained using a homotopic erosion technique [Mangin et al., 1995]. A "crevasse detector" is used to reconstruct sulcal structures as the medial surfaces from two opposing gyral banks that span from the most internal point of the sulcal fold to the convex hull of the cortex [Mangin et al., 1995, 2004] (Fig. 1E). The BrainVISA sulcal identification pipeline incorporates a congregation of 500 artificial neural network-based pattern classifiers. Each classifier is tuned to identify a particular feature of the sulcal patterns. The neural networks work collaboratively to identify 56 sulcal structures for each hemisphere. The networks were trained on a database of 26 expertly classified images. Classification

accuracy ranged from 95% for primary and secondary fissures to around 70% for the highly variable tertiary sulcal structures [Rivière et al., 2002]. Two skilled neuroanatomist reviewed the results of the automated sulcal identification. In an infrequent event (~5–15%) where a sulcus was assigned an incorrect label, the sulcus was manually relabeled while overlaid on the cortical surface (Fig. 1F). Special attention was paid to anastomotic sulci, small, accessory sulci, usually less than 5 mm in length [Ono et al., 1990]. Anastomotic sulci were excluded from analysis because a span measurement performed at intersections between anastomotic and targeted sulci, especially along the direction of the anastomotic sulcus, would lead to exaggerated values.

Five cortical regions were composed from twelve cortical sulci by grouping functionally similar sulci (Table I). Sulci selected for this analysis were previously used to study age-related changes during senescence [Kochunov et al., 2005b]. All sulci have very high inter-subject incidence rates (~99%) [Ono M et al., 1990] and were present in all subjects.

**FA processing**—We used the tract-based spatial statistics (TBSS) software [Smith et al., 2006] for multi-subject analysis of FA [Kochunov et al., 2007]. TBSS combines strengths of voxelwise [e.g., VBM—voxel based morphometry [Ashburner and Friston, 2000] and WM tractography methods. During TBSS processing, FA images were created by fitting the diffusion tensor to the raw diffusion data [Smith, 2002]. Then, all FA images were nonlinearly aligned to a group-wise, minimal-deformation target (MDT) brain. The group's MDT brain was identified by warping all individual brain images in the group to each other [Kochunov et al., 2001]. Individual FA images were averaged to produce a group-average anisotropy image. This image is used to create a group-wise skeleton of white matter tracts. Finally, FA images were projected onto the group-wise skeleton of white matter structures. This step accounted for residual misalignment among individual white matter tracts. In the last step, we used a threshold FA value of 0.2 to reconstruct fully connected skeleton for even the eldest subjects where FA values were generally lower.

**HWM processing**—Hyperintense white matter (HWM) volume was measured from T2weighted FLAIR images as described by Kochunov et al., [2007]. FLAIR images were preprocessed by removal of nonbrain tissue, registration to the T1-weighted images/ Talairach frame, and RF inhomogeneity correction.

## Structural Indices of Cerebral Integrity

Descriptions of methods for measurements of global and regional indices of cerebral integrity were published elsewhere [Kochunov et al., 2005b, 2007,2008). Cortical cerebral integrity measurements, sulcal, intergyral span, and GM thickness were performed for five regions (Table I); FA values were measured for the entire brain and three regions of corpus callosum; HWM volume was calculated for the entire brain.

**Sulcal span**—Sulcal span was defined as the average span of the sulcal space along the normal projections to the medial sulcal mesh (see Fig. 2). High resolution medial sulcal meshes for each sulcus were created using the BrainVISA sulcal extraction pipeline. Geometrically, a medial sulcal mesh traverses the sulcal space in the middle of the sulcal "span" dimension, parallel to the gyral gray matter borders, and spans the entire sulcal "depth" dimension (see Fig. 2). The medial sulcal mesh for each sulcus was used to seed a 3D region growing algorithm to mask cerebrospinal fluid volume associated with the sulcal structure (see Fig. 2). To prevent cerebrospinal fluid from "spilling over" into intersecting sulci, points that were further than 10 mm away from the sulcal skeleton were rejected. The convex hull of the cortical gray matter ribbon was then used to isolate sulcal space from subarachnoid cerebrospinal fluid. Sulcal span was calculated as the Euclidean distance between two points on the gyral gray matter mesh on

either side of the sulcal surface (see Fig. 2). The point of intersection between the normal projection and the gray matter mesh corresponded to voxels with partial cerebrospinal fluid/ gray matter values of 50%.

**Intergyral span**—Intergyral span was calculated by extending the sulcal span tracings until they intersected with the surface corresponding to the gyral gray matter-white matter interface. Gyral span was defined as the Euclidean distance between two points residing on the gyral white matter mesh on either side of the sulcal surface (see Fig. 2).

**Gray matter thickness**—Gray matter thickness was calculated by starting a new tracing at the point of intersection between a normal projection to the sulcal mesh and the exterior gray matter mesh. To measure GM thickness, a normal projection to the exterior gray matter mesh was traced until it intersected with the interior gray matter mesh that describes the gray matter-white matter interface (see Fig. 2). The point of intersection between the normal projection and the interior gray matter mesh corresponds to voxels with partial white matter/gray matter values of ~50%.

**Fractional anisotropy**—FA values were measured as the whole brain average and for three regions of the corpus callosum (CC) subdivided along its anterior-posterior length [Kochunov et al., 2005a]. The whole brain FA average value was calculated for all voxels of the skeleton. Regional CC FA values were calculated from a 10 mm wide band of median CC. This band was subdivided along the A-P direction into three equal subparts: Genu, midbody. and splenium. Studies in cadavers indicated that the genu (anterior third) contains the thinly myelinated, tightly packed association fibers connecting the bilateral prefrontal cortices; the midbody (middle third) primarily contains thickly myelinated fibers for motor, somatosensory, and auditory cortices; and the splenium (posterior third) contains a mixture of heavily myelinated and thinly myelinated fibers connecting temporal, parietal, and occipital lobes [Aboitiz, 1992; Aboitiz et al., 1992; Highley et al., 1999].

**HWM volume**—Regions of HWM were manually delineated (painted) on the FLAIR images by two experienced neuroanatomists with high (r = 0.85) inter-rater reliability. HWM volume was calculated by adding volumes of individual regions.

#### Statistic analysis

Normality of EXIT scores and cerebral integrity indices was tested by Shapiro-Wilk's and Kolmogorov-Smirnov tests. Logarithmic transformation was used to achieve normality for the EXIT scores and HWM volume. Linear and age-corrected partial correlation analysis was used to examine relations among indices of structural integrity and normalized EXIT Scores (NES). Only the indices that showed significant age-corrected correlation with NES were selected for the further analysis. This selection criterion allowed us to study age-independent association between structural indices and cognition. It also allowed for the reduction of the number of dependent variables prior to factor analysis to achieve a ratio of dependent variables to the number of subjects ratio of at least 1:10 [Osborne and Costello, 2004]. Factor analysis was performed to further reduce the number of dependent variables and to remove the colinearity among the indices of cerebral integrity to satisfy the requirements of regression analysis. Factor analysis used the principal components analysis (PCA) to obtain linear composites of correlated variables. It yielded eigenvector values describing correlations between a variable and a component. A varimax rotation was then used to remove colinearity e.g., to orthogonize the individual eigenvectors. Results of factor analysis yielded factor loadings (correlations between a variable and a factor) and factor scores (a subject's standardized score on each factor). Finally, a two-stage regression was performed to probe the multivariate effects of age and structural indices of cerebral integrity on the inter-subject variability in NES. Subject age

was entered at first, followed by the factor scores extracted from structural measurements. The regression analysis yielded the degree of variance described at each entry step and whether the change was significant. It also produced standardized coefficients ( $\beta$ ) that estimate linear associations between criterion variables (NES) and predictor variables (factor scores extracted from the structural indices).

## RESULTS

## Normalized EXIT Scores

In the maturing group, raw EXIT scores ranged from 3 to 16 (average =  $7.5 \pm 2.7$ ); in the senescing group raw EXIT scores ranged 5–25 (average =  $12.8 \pm 5.0$ ). EXIT scores were adjusted using logarithmic transformation to achieve normality, producing normalized EXIT scores (NES) that satisfied this criteria (maturing group: Shapiro-Wilk's w = 0.95, P = 0.13; Kolmogorov-Smirnov F = 0.13, P = 0.19; senescing group: Shapiro-Wilk's w = 0.94, P = 0.13; Kolmogorov-Smirnov F = 0.13, P = 0.15).

Average NES values were significantly different between groups  $(1.96 \pm 0.06 \text{ maturing group}; 2.5 \pm 0.07 \text{ senescing group}; t = 5.5; P < 1e-5)$ . In addition, age-related slopes for NES were opposite in sign between the maturing and senescing groups (see Fig. 3), and the between-group difference in correlation coefficients with age was highly significant (Fisher's z = 3.7; P < 0.001). Younger subjects showed a nonsignificant negative correlation between NES and age, hinting that performance on the test improved with age. NES were positively and significantly correlated with age in the senescing group; tracking an expected decline of ECF with age (Table II; Fig. 3).

## Whole-Brain Indices of Cerebral Integrity

Whole-brain average measurements of cerebral integrity (GM thickness, intergyral and sulcal spans, FA, and HWM volume) were significantly correlated with age in the senescing group (Table II; Fig. 4A–C; Fig. 5B). In the maturing group, only the FA measurements in the genu of CC showed a statistically significant correlation with age (Figure 4A–C, Figure 5A). Partial correlation analysis was used to probe age-independent trends between NES and whole-brain average measurements. The only significant age-independent correlation was between NES and the whole brain average sulcal span in the senescing group (Table III).

## **Regional Indices of Cerebral Integrity**

The results of uncorrected and age-corrected correlation analyses between NES and three indices of cerebral integrity (GM thickness, intergyral, and sulcal spans) for five cortical regions for two hemispheres are presented in Table IV and Table V. The level of statistical significance was set at P = 0.001 to reduce the probability of Type 1 errors associated with 30 (three measurements \* five regions \* two hemispheres) comparisons.

In the maturing group, none of the 30 cortical integrity measurement showed a significant (P < 0.001) correlation with NES. However, there were marginally significant correlations ( $P \sim 0.01$ ) observed between NES and right and left sulcal span in the occipital region (Table IV).

In the senescing group, 13 of 30 cortical integrity measurements were significantly (P < 0.001) correlated with NES (Table V). Age-corrected correlation analysis revealed three regional measurements that correlated with NES at P < 0.001 (left and right sulcal span and right hemispheric integral span in the superior frontal/anterior cingulate region).

#### **Factor Analysis**

Factor analysis was performed in senescing group for regional cortical indices that showed a significant (P < 0.001) age-corrected partial correlation with NES (Table VI). Factor 1 loaded on left and right sulcal span measurements for the superior frontal/anterior cingulate sulcus. It was significantly and positively correlated with NES, pr = 0.61; P < 0.001. Factor 2 loaded on intergyral span measures for the right superior frontal/anterior cingulate sulci. It was also significantly and positively correlated with the NES, pr = 0.57, P < 0.001.

## **Regression Analysis**

A two-stage regression analysis was performed in the senescing group. NES was used as the dependent variable and was regressed versus age and the two factor scores. In the first stage, age was entered as the only independent variable. In the second stage, factor scores were entered as independent variables using a step-wise procedure. Age and factor scores produced statistically significant changes in r ( $\Delta r \ge 0.33$ ;  $P \le 0.01$ ). However, the regression coefficient ( $\beta$ ) for age was not significant, indicating that factor scores extracted from structural measurements fully captured age-related variability in NES (Table VII).

## DISCUSSION

This study sought to explore the relationship between cognition and indices of cerebral integrity in normally aging subjects. Our results support a multimodal MRI approach to help predict cognitive changes in normal aging. As noted in the introduction, Royall et al., [2000] have previously shown a inverse "U" relationship between the EXIT and age, that peaks in the 3rd to 4th decade of life [Royall, 2000]. In the current study, trends in the EXIT scores showed that ECF improved with age in the younger, maturing group and declined in the older, senescing group, thereby replicating the previous work. Parallel to ECF trends, indices of cerebral integrity hinted at developmental trends in the maturing group and showed an age-related decline in the senescing group.

In the maturing group, age-related trends in NES and indices of cerebral integrity hinted at ongoing cerebral development, possibly because of late myelination of associative white matter tracts. This was exemplified by the increasing FA values in the genu of the CC, which is composed of thinly myelinated, densely packed, associative commissural fibers that link prefrontal and frontal areas [Aboitiz, 1992; Aboitiz et al., 1992]. An age-related increase in the genu's FA values in maturing subjects was reported to be an indication that the associative WM tracts continue to develop into the third decade of life [Abe et al., 2002; Huppi et al., 1998; Sullivan et al., 2001]. In contrast, FA values in the body of CC were not significantly correlated with age (Table II). This was expected as the fiber tracts located in the body of CC are composed of heavily myelinated motor and sensory fibers, known to myelinate at an early age [Kochunov et al., 2005a]. We also believe that the observed, albeit not statistically significant, age-related decrease of the intergyral span is further evidence of late myelination. We tested this observation by analyzing the correlation between the intergyral span and FA measurements in the genu of CC, which was highly significant (r = -0.55; P < 1e-3). In contrast, the correlation between intergyral span and FA values in the body of CC was not significant (r = -0.12; P = 0.5). These results suggest that NES and indices of cerebral integrity were sensitive to cerebral maturation in the younger group.

In the senescing group, age-related trends in NES and indices of cerebral integrity indicated a decline in cognition with a corresponding decline in cerebral integrity. NES and all cerebral indices showed statistically significant (P < 0.05) correlation with age (Table II). The highest correlation with age was observed for sulcal span (r = 0.68), followed by FA in the genu of the CC (r = -0.63) and GM thickness (r = -0.61). The smallest correlations were seen for the

HWM volume and FA measurements for the body of CC (Table II). Interestingly, FA in the genu of CC was the only cerebral index that showed statistically significant association with age in both maturing and senescing groups, suggesting that FA of late-myelinated associative fibers was sensitive to both maturation and senescence.

In the maturing group, regional measures of cerebral integrity were poor at predicting changes in NES (Table IV). We found marginally significant correlation with NES for the sulcal span of occipital areas (age-corrected, pr = 0.38;  $P \sim 0.01$ ) (Table IV; Fig. 6). However, these structures (e.g. left and right occipital structures) were previously reported to have age-related increases in cortical complexity in children ages 6-16 [Blanton et al., 2001]. Blanton et al. hypothesized that changes in axonal tensions due to continuous myelination of associative fibers were the likely cause [Blanton et al., 2001]. We tested their hypothesis by evaluating age-controlled correlation between occipital sulcal span and FA values in the CC. The strongest correlation was observed with the FA of the splenium of the CC (age corrected, pr = -0.44, P < 0.01). The splenium of CC contains late myelinating, associative visual-spatial orientation fibers linking occipital and parietal lobes [Aboitiz et al., 1992]. Occipital sulcal span measures also significantly correlated with FA values in the other late-myelinating area of CC, the genu (r = -0.37, P < 0.05), but were not correlated with FA in the mostly early-myelinated of CC midbody (r = -0.29;  $P \sim 0.10$ ). This confirms that maturational changes in occipital areas, first reported by Blanton et al., [2001], are likely due to late myelination of associative WM. Thus, marginally significant correlation between occipital sulcal span and NES is indirect evidence that late-term maturation of cerebral WM influenced variability in ECF in maturing group.

In the senescing group, regional measures of cerebral integrity correctly localized the ECF network and identified specific neuroanatomical correlates predictive of decline in NES (Table IV). Regional measurements of sulcal and intergyral span from frontal and anterior-cingulate areas accounted for 62% of the intersubject variability in NES. Localization of the ECF network to the frontal-cingulate areas is consistent with previous reports from our center [Royall et al., 2001] and others [Collette et al., 2005,2006;Duncan and Owen, 2000;Vandenberghe et al., 2000]. An important finding was that only sulcal and intergyral spans in the anterior cingulate and superior frontal areas were associated with age-independent decline in NES. Factor analysis distilled these measurements into two orthogonal categories each explaining equal amounts of variance in NES (Table VI). Factor 1 was significantly correlated with two common measures of WM atrophy: The volume of HWM (age-corrected, pr = 0.46; P < 0.01) (see Fig. 7] and whole-brain average FA value (age-corrected, pr = -0.45; P < 0.01). These associations were expected, as increases in HWM volume are frequently reported to be associated with cognitive decline in normal aging individuals [Gunning-Dixon and Raz, 2000,2003;Raz et al., 1998]. Factor 2, was significantly correlated with FA values of the genu of the CC (age corrected, pr = 0.34; P < 0.05) (see Fig. 8). This is a novel finding suggesting that interhemispheric connections through the genu are associated with a decline in NES.

#### Limitations

A possible limitation of this study was the use of EXIT as the measure of ECF. ECF is a multidimensional construct comprising a variety of cognitive processes. This would raise the question as to whether the EXIT findings are nonspecific projections from multiple dimensions. However, previous research done by this group and others has shown that the EXIT loads on a specific ECF dimension that is uniquely associated with disability and is not colabeled by either the WCST or trail-making [Park et al., 2006] tests (TMT) [Lewis and Miller, 2007; Royall et al., 2003]. It has also been shown that EXIT scores are closely related to the Tower of London (TOL) and similar cognitive planning tasks strongly associated with measures of functional status [Lewis and Miller, 2007; Royall et al., 1992]. The association between planning and functional status helps to explain the EXIT's singularly strong associations with

functional outcomes [Lewis and Miller, 2007]; particularly relative to the WCST or TMT (i.e., putative ECF measures that do not co-label the same ECF factor as the EXIT) and are less strongly associated with functional status [Bell-McGinty et al., 2002]. Another limitation of this investigation was the relatively narrow age range of the maturing, younger group. This restriction could have contributed to the lack of significant findings for this group through a variance compression artifact. In particular, younger subjects had smaller variability in age and NES versus the older group (SD = 2.7 vs 5.0).

Finally, it is important to consider the statistical approach taken here. We conducted the analyses with categorization of the subjects into two groups. Ideally, this analysis should have been performed by entering subject's age as a continuous variable. Then, a nonlinear regression/correlation analysis could have been used to model the effects of the age as a quadratic or higher-order function. However, the relatively low number of subjects and sparseness of the data in the age range of 35–55 years prevented us from performing a nonlinear analysis. In addition, while subjects from maturing and senescing age groups clearly showed diverging age-related trends for neuropsychological and neuroanatomical measurements (e.g. Figure 4 and Figure 5), these trends have clear linear, rather than higher-order components.

## CONCLUSION

We analyzed an association between neuropsychological assessment of ECF and regional indices of cerebral integrity in healthy maturing and sensing populations. In the maturing population of 19–29 years old, improvements in ECF corresponded to cerebral maturation trends in the indices cerebral integrity. In the senescing population of 30–90 years old, decline in ECF corresponded to trends in neuroanatomical decline of the indices of cerebral integrity. In both populations, whole-brain trends average cerebral indices did not strongly correlate with trends in ECF. However, regional trends in cerebral markers correlated strongly with trends in ECF. In maturing subjects, the regional pattern of correlation with ECF hinted at late myelination of associative tracks of cerebral WM. In older subjects, regional association patterns included fronto-cingulate structures that were previously reported to be involved in the modulation of ECF. In summary, regional indices of cerebral integrity, corresponding to atrophy of cerebral WM, were highly predictive of deficits in ECF.

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#### Figure 1.

T1-weighted image processing pipeline for extraction of indices of cortical integrity consists of skull stripping (**A**); spatial normalization, radio-frequency homogeneity correction, and tissue segmentation (**B**); extraction of gray and white matter surfaces (**C**,**D**) extraction of sulcal folds (**E**); and automated labeling of sulci (**F**).



#### Figure 2.

Three indices of cortical integrity were measured for primary cortical sulci (Table I). The sulcal surface is defined as a surface that spans the median space between two gyral banks. Three measurements are computer per sulcal structure: Average sulcal and intergyral spans. A sulcal span (solid line) was defined as a 3D distance between opposing points on the GM mesh along the normal projections to the medial sulcal mesh. A intergyral span (dotted line) was defined as the 3D distance between opposing points on the WM mesh along the normal projection to the medial sulcal mesh. These measurements are performed for all nodes of the individual sulcal mesh, resulting in averaging of ~2–5000 measurements for each sulcal structure.

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#### Figure 3.

Age-related trends for Executive Interview scores for maturing ( $\circ$ , dotted line), age 19–29 and senescing ( $\bullet$ , solid line), age 30–90 years subjects.

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#### Figure 4.

Age-related trends for average cortical indices of cerebral integrity: GM thickness (**A**), WM span (**B**), and Sulcal Span (**C**) for the maturing ( $\circ$ ), age 19–29 and senescing ( $\bullet$ ), age 30–90 years subjects.

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Figure 5.

Age-related trends for average subcortical indices of cerebral integrity: Average fractional anisotropy of water diffusion (A) and volume of hyperintense white matte for the maturing ( $\circ$ ), age 19–29 and senescing ( $\bullet$ ), age 30–90 years groups of subjects.



#### Figure 6.

Age-corrected normalized Executive Interview scores (NES) plotted versus bilateral measurements of sulcal span in occipital region for maturing subjects (age 19–29).

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## Figure 7.

Factor 1 scores in senescing subjects (ages 30–90) are plotted vs. hyperintense white matter (HWM) volume.



## Figure 8.

Average fractional anisotropy values for the genu, shown as the shaded area of the corpus callosum, are plotted versus factor scores for Factor 2 in senescing subjects (age 30–90)

#### TABLE I

Gyral gray matter thickness, sulcal and gyral spans were calculated for five cortical regions per hemisphere based on 12 primary and secondary sulcal structures per hemisphere

Region	Sulcal structures and brodmann areas (BA)
Superior frontal/anterior cingulate	Superior frontal sulcus (BA 46/9/8); anterior cingulate sulcus(BA 24/32)
Primary somatosensory	Pre-central sulcus; postcentral sulcus; central sulcus (BA 6/4/3/1/2)
Parietal/post.cingulate	Intraparietal fissure (BA 19/39); posterior cingulate sulcus (BA 23/31)
Temporal	Superior temporal sulcus (BA22/21); collateral sulci(BA20/36)
Occipital	Occipital/calcarine sulci(BA 17/18); parietaloccipital fissure(BA 7/19)

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Linear correlation coefficients (r) between subject's age and normalized executive interview scores (NES); and whole-brain indices of TABLE II cerebral integrity

Correlation with age	NES	GM thickness	WM span	Sulcal span	FA	FA genu	FA body	FA splenium	HWM volume
Maturing group Senescing group	-0.21	-0.11	-0.25	-0.19	0.15	$0.40^*$	0.19	0.31	N/A
Senescing group	0.60*	-0.61	0.49*	$0.68^*$	-0.53*	-0.63	$-0.38^{*}$	-0.51*	.37*
HWM volume could not be	measured for	younger subjects.							

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. Indicated statistically significant (P < 0.05) correlation coefficients.

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 TABLE III

 Partial, age-corrected correlation coefficients (pr) between normalized executive interview scores (NES) and indices of cerebral integrity

Correlation with NES	GM thickness	WM span	Sulcal span	FA	FA genu	FA body	FA splenium	HWM volume
Maturing group	0.06	0.15	0.31	-0.17	-0.08	-0.14	-0.17	N/A
Senescing group	-0.04	0.22	$0.36^*$	-0.23	-0.21	0.03	-0.20	0.27

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HWM volume could not be measured for younger subjects.

\* Indicated statistically significant (P < 0.05) correlation coefficients.

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Uncorrected and age-corrected partial correlation between regional, right (R)/left (L) cerebral integrity indices and normalized executive interview scores for the younger, maturing (age range 19-29, N = 34) subjects **TABLE IV** 

	R. GM tl	hickness	L. GM th	iickness	R. WM	1 span	L. WN	A span	R. sulc	al span	L. sulc	al span
Raw and age-corrected partial correlation with NES	Direct	Partia	Direct	Partial	Direct	Partial	Direct	Partial	Direct	Partial	Direct	Partial
S. Frontal/anterior cingulated	-0.05	-0.05	0.04	0.00	0.04	0.00	0.06	-0.10	0.21	0.18	0.19	0.13
Primary somatosensory	-0.02	-0.05	-0.03	-0.03	-0.02	-0.02	0.18	-0.04	0.18	0.16	0.25	0.22
Parietal/post cingulate	0.03	-0.05	0.06	-0.03	0.17	0.08	0.20	0.17	0.26	0.23	0.19	0.18
Temporal	-0.23	0.11	0.05	-0.03	0.24	0.12	0.14	0.17	0.31	0.30	0.05	0.01
Occipital	-0.35	-0.34	-0.14	-0.15	-0.02	0.06	0.17	0.11	$0.42^*$	$0.38^*$	$0.41^*$	$0.38^*$
Average	-0.13	-0.12	0.02	-0.05	0.08	0.05	0.15	0.06	0.28	0.25	0.22	0.18
*												

Indicated statistically significant (P < 0.05) correlation coefficients.

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Uncorrected and age-corrected partial correlation between regional, right (R)/left (L) cerebral integrity indices and normalized executive interview scores (NES) for older (O), senescing (age-range 30-90; N = 38) subjects **TABLE V** 

	R. GM tł	nickness	L. GM th	ickness	R. WN	1 span	T. WN	1 span	R. sulc	al span	L. sulca	ıl span
Raw and age-corrected partial correlation with NES	Direct	Partial	Direct	Partial	Direct	Partial	Direct	Partial	Direct	Partial	Direct	Partial
S. Frontal/anterior cingulate	-0.34	-0.08	-0.33*	-0.05	0.68*	0.58*	$0.56^*$	$0.33^*$	0.67*	$0.61^*$	$0.62^{*}$	$0.54^{*}$
Primary somatosensory	$-0.41^{*}$	-0.11	$-0.35^{*}$	-0.09	0.22	0.03	0.31	0.02	$0.57^*$	0.26	$0.55^*$	0.26
Parietal/post Cingulate	$-0.37^{*}$	-0.09	$-0.32^{*}$	-0.14	0.17	0.02	0.20	-0.08	$0.40^*$	-0.02	$0.40^*$	0.11
Temporal	$-0.39^{*}$	-0.25	$-0.36^{*}$	-0.15	0.09	-0.05	0.16	-0.16	0.55*	0.34	$0.40^*$	0.13
Occipital	$-0.35^{*}$	0.03	$-0.48^{*}$	-0.11	0.03	0.03	0.04	-0.08	0.27	0.11	$0.35^*$	0.11
Average	-0.37	-0.1	-0.36	-0.11	0.23	0.11	0.25	-0.08	0.49	0.26	0.46	0.23
Bolded numbers indicate statistica	Ily significant of	correlations at I	<sup>p</sup> < 0.001 level									

\* Indicated statistically significant (P < 0.05) correlation coefficients.

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## TABLE VI

## Factor analysis in senescing group

	Factor 1 (60%)	Factor 2 (37%)
R. Sup. Frontal/A. Cingulate sulcal span	0.93	0.36
L. Sup. Frontal/A. Cingulate sulcal span	0.93	0.35
R. Sup. Frontal/A. Cingulate gyral wm span	0.36	0.94

ression result in the senescing group	Adjusted $\Delta r$ Significance Standardized Significance $r$ of $\Delta r$ $\beta$ of $\beta$	0.58 0.60 <0.01 0.00 0.12	0.66 0.33 =0.01 0.19 0.001	0.77 0.35 <0.01 0.18 0.002	
Iwo-stage regression result in the senesci	r Adjusted	0.60 0.58	0.68 0.66	0.79 0.77	
Ľ	Model	Age	Factor 1	Factor 2	

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