Cerebral correlates of cognitive aging: Gray-white-matter differentiation in the medial temporal lobes, and fluid versus crystallized abilities

NAFTALI RAZ
Memphis State University, Memphis, Tennessee

DARYL MILLMAN
Chicago Medical School, North Chicago, Illinois

and

GÜNSELİ SARPEL
North Chicago VA Medical Center, North Chicago, Illinois

We investigated the relationship between age, structural properties of selected cerebral regions, and cognitive performance in healthy adults, 18 to 78 years old. Spin-lattice relaxation time (T1), measured by nuclear magnetic resonance, was used to describe the structural composition of the brain tissue. Temporal lobe white-matter T1 showed age-related prolongation best described by a quadratic polynomial. There was a significant cubic trend in the association of hippocampal (gray-matter) T1 with age. In the examined regions of the medial temporal lobes, normally observed differentiation between gray- and white-matter T1 diminished linearly with age and disappeared almost completely in the elderly. Age and the ratio of gray- to white-matter T1 accounted for 53% of the variance in a measure of fluid intelligence (Cattell Culture Fair Test); the unique contributions of age and of gray-white-matter T1 ratio were 23% and 3%, respectively. The largest share of the variance in fluid intelligence (27%) was explained by the common influence of age and gray-white-matter T1 ratio. The same set of variables explained no significant proportion of the variance in crystallized intelligence. The possible mechanisms underlying age-related changes in gray-white-matter differentiation, their relationship to age-related selective deterioration of cognitive functions, and the implications of the findings for research on biological markers of aging are discussed.

Aging is associated with specific changes in cerebral morphology, as well as with differential decline in cognitive abilities (Coleman & Flood, 1987; Horn, 1986). Cumulative evidence from postmortem studies suggests that the regions of the brain that are more likely to undergo age-associated alterations are the medial temporal lobes encompassing the hippocampal formation and the prefrontal cortex (Barnes, 1983; de Leon, George, Stylopoulos, Smith, & Miller, 1989; Mani, Lohr, & Jeste, 1986; Terry, DeTeresa, & Hansen, 1987). In the cognitive domain, a group of abilities such as nonverbal reasoning, rule discovery, and concept formation, usually called fluid intelligence, is especially vulnerable to the effects of aging. On the other hand, formal verbal reasoning, comprehension of culture-specific rules and strategies, and general fund of knowledge—the abilities constituting crystallized intelligence—do not decline with age (Horn, 1986).

Pathological changes in the temporal lobes, mainly in the hippocampus, have been implicated in cognitive declines observed in the normal elderly and are regarded as one of the major signs of age-related cognitive pathology (Squire, 1986; Winocur, 1988). Horn (1986) has suggested that cognitive components constituting fluid intelligence can be affected by lesions in the medial temporal and limbic structures. He also reported age-related decrease of regional cerebral blood flow (rCBF) in broadly defined medial temporal aspects of the brain. A relationship between the extent of hippocampal atrophy and memory deficit has been demonstrated in several cases of traumatic amnesia (Press, Amaral, & Squire, 1989).

Until recently, a direct test of any hypothesis relating differential decline of cognitive abilities to age-related deterioration of cerebral structures has been impossible due...
to the lack of instruments permitting an in vivo assessment of neuroanatomy and brain function in healthy individuals. Introduction of computerized imaging techniques—computer-assisted tomography (CT) and magnetic resonance imaging (MRI)—has created new opportunities for evaluation of age-related neuroanatomical changes in intact humans. In addition to allowing visualization of brain structures in any anatomical plane, MRI provides an opportunity for an in vivo evaluation of the molecular behavior of water, the main constituent of cerebral tissue (Harms & Kramer, 1985).

Contemporary MRI scanners provide two basic parameters describing the behavior of water protons in living tissue: spin-lattice (longitudinal) relaxation time (T1) and spin-spin (transverse) relaxation time (T2). Both T1 and T2 are sensitive to differences in molecular organization of healthy and diseased tissue (Bottomley, Foster, Argersinger, & Pfeifer, 1984). Although for a given magnetic field strength T1 correlates highly with water content of the brain \( r > .9 \), it is influenced by changes in the macromolecular structure of water as well (Bottomley et al., 1984; MacDonald et al., 1985; Unger, Littlefield, & Gado, 1988). The dependence of T1 on the relative share of bulk-phase and hydration-layer water makes it a sensitive tissue-typing tool (Mathur-DeVre, 1984). In general, tissue containing a substantial amount of bulk water has longer T1 than tissue in which water is represented, mainly in hydration layer clinging to macromolecules (Mathur-DeVre, 1984; Unger et al., 1988). As a result, cerebral gray matter would normally have considerably longer T1 time constants than the white matter.

Water content of the white matter increases dramatically with age (Wiggins et al., 1988). Focal edema due to microinfarctions (Tomonaga, Yamanouchi, Tohgi, & Kameyama, 1982), atrophic perivascular demyelination (Ansari & Loch, 1975), cumulative hypoxic-ischemic damage (Ginsberg, Hedley-Whyte, & Richardson, 1975), and loss of proteins and glycolipids from the myelin sheath (Kirkpatrick & Hayman, 1987; Wiggins et al., 1988) are associated with aging and may produce molecular changes causing prolongation of T1 in the white matter. Indeed, perivascular demyelination has been histopathologically confirmed in many cases of the white-matter lesions detected with MRI (Kirkpatrick & Hayman, 1987). Wahlund et al. (1990) reported that age-related changes in white-matter composition are reflected in altered T1 values. In their study, conducted on an age-restricted sample of elderly subjects, the correlation between age and frontal lobe white-matter T1 averaged across the hemispheres was \( r = .60 \). In a recent study of a representative sample of normal volunteers, Jernigan et al. (in press) reported a multiple correlation of \( r = .68 \) between a quadratic function of age and white-matter T2.

The T1 values of the gray matter may also be affected by a variety of age-related processes. Postmortem studies indicate that aging is accompanied by a decrease in ne-
employed. All subjects scored at least 29 out of 30 points on the Mini-Mental Status Examination (MMSE; Folstein, Folstein, & McHugh, 1975). The mean age in this sample, which included 15 males and 11 females, was 45.8 years ($SD = 21.8$), with a range of 18 to 78 years. The average duration of formal education was 14.7 years; the correlation between age and education was $r = - .29$, n.s. No formal assessment of handedness was conducted, but all subjects were self-reported right-handers. They signed an informed consent form and were thoroughly briefed regarding the objectives of the study.

**Apparatus and Procedure**

Imaging and spin-lattice relaxation time (T1) measurements were performed on a Fonar Beta-3000.3T permanent-magnet scanner at the McCormick University Clinics (University of Health Sciences/Chicago Medical school). All imaging was conducted in a spin-echo (SE) mode with a $256 \times 256$ matrix and a 22-cm field of view. A 27-cm head coil was used in all procedures.

Before the T1 measures were taken, two imaging sequences were performed. Nine sagittal slices were obtained at echo time (TE)/repetition time (TR) = 28/600 msec, with three averages per projection and slice thickness/interval of 4.2/6.0 mm. Upon completion of the 7-min sagittal sequence, a coronal sequence was initiated. In that sequence, 17 to 21 slices (depending on the cranial size) were taken in the coronal plane, with TE/TR = 28/1,800 msec; two averages per projection, and slice thickness/interval of 6.6/8.6 mm. The midsagittal slice was used as a scout, and the position of the coronal slices was determined in reference to the middle of the pituitary gland. Acquisition of the coronal sequence took about 15 min.

All scans were examined by a neurologist with extensive experience in clinical MRI (G.S.). One subject (76-year-old female) showed moderate parietal atrophy, and 2 subjects (71-year-old male

![Figure 1. Top: A coronal MRI slice through anterior hippocampal formation used for T1 measurement. Bottom: The same slice with superimposed T1 values.](image-url)
and 68-year-old female) had minor periventricular patches of increased signal intensity. Otherwise, no space-occupying lesions or significant areas of signal hyperintensity were observed, and scan appearance was judged normal for age. As indicated by the MMSE scores and the interview, the subjects with positive neuroradiological findings exhibited neither clinical–behavioral harbinger of mental deterioration nor a pattern of risk factors characteristic of demening illness. The finding of areas of hyperintensity per se is not indicative of pathology (Leys et al., 1990; Sze et al., 1986) and does not necessitate exclusion of the subjects.

Measurement of T1

The measurements were conducted on four regions of interest (ROI) on a single slice. After the imaging sequences were completed, the subject remained in his/her position, and a coronal slice (Figure 1, top) was selected to obtain the T1 values. That slice—third caudally from the middle of the pituitary—was selected because it allowed a clear view of the hippocampal formation. The selection was verified against the atlas of correlative MRI neuroanatomy (Daniels, Naughton, & Naidich, 1987). Although the investigators who performed T1 measurement were not blind to the subjects' age, the use of clearly defined neuroanatomical landmarks and uniform slice selection procedures made bias in selection of the ROI rather unlikely.

The T1 values were computed for the anterior part of the hippocampal formation and the adjacent subcortical white matter in both hemispheres. The computations were performed on-line using a strip method developed by the Fair Corporation, Melville, NY, and described by Morone et al. (1987). In this method, the acquisition of T1 is separated from the imaging process. It is performed with an SE pulse sequence and time-varying gradients at 13 different repetition rates. At the shortest repetition time (TR), signal intensity is the greatest, dropping exponentially with prolongation of the TR. The T1 measurement is based on the linear regression of the logarithm of relative decrement in signal amplitude on the repetition time. In comparison with conventional methods, in which T1 is computed off-line and is based on only two TR values, the strip method is distinguished by relatively high reliability. Its main drawback is that the measurement is restricted to one relatively small region at a time.

Thirteen TR values were used in each T1 strip computation, and the total acquisition time was 3.5 min. The spatial resolution of the T1 measurement was 7 mm, and, for each strip, T1 values from up to 10 points along the strip were recorded. An example of a strip with 10 T1 values displayed is presented in the bottom panel of Figure 1. The T1 measurement sequence was repeated until at least six measurements per ROI were obtained. The median T1 value for a given ROI served as its representative T1.

Psychometric Tests

To assess cognitive functions, two tests, one representative of fluid and one of crystallized intelligence, were used. The Cattell Culture Fair Test (CFIT; Cattell & Cattell, 1973) and Extended Vocabulary (V3 from Educational Testing Services Factor-Referenced Test Kit (Eckstrom, French, Harman, & Derman, 1976) were administered to each subject individually in a span of no more than a week from the date of MRI scanning. The total number of correct responses on CFIT was used as a measure of fluid intelligence. The V3 score used as a measure of crystallized intelligence was computed by subtracting the number of incorrect answers divided by four from the total number of correct answers, in order to adjust for guessing.

RESULTS

To reduce the number of variables in a design with few degrees of freedom, T1 values for medial temporal gray and white matter were averaged across the hemispheres. Zero-order correlations among the cognitive, demographic, and radiological variables are presented in Table 1.

As hypothesized, the white-matter T1 exhibited a positive linear relationship with age. Because of Jernigan et al.’s (in press) finding of nonlinearity in age–T2 relationship, the data were examined for presence of higher order trends. The polynomial regression analysis revealed a significant quadratic trend that significantly increased the correlation between the white-matter T1 and age to \( r = .63, p < .005 \). The best-fit quadratic equation describing this relationship was: \( T1_w = 622.33 - 4.68 \cdot \text{Age} + .066 \cdot \text{Age}^2 \).

The relationship between the gray-matter T1 and age was more complex. The linear correlation between the two variables was negative, as predicted, but small and statistically nonsignificant. However, the polynomial regression analysis revealed a significant cubic trend that increased the multiple correlation between the gray-matter T1 and the age components to \( r = .54, p < .05 \). The relationship between the hippocampal T1 and age was described by a cubic equation: \( T1_h = 394.98 + 20.98 \cdot \text{Age} - .528 \cdot \text{Age}^2 + .004 \cdot \text{Age}^3 \). No correlation between gray- and white-matter T1 was found.

The percent ratio of gray- to white-matter T1 was linearly related to age, approximating 100\% in the older subjects. As evident from the distribution in Figure 2a, the negative association between age and gray–white-matter differentiation index in the medial temporal lobes was not disproportionately influenced by outliers. Nonlinear regression analysis revealed no significant higher order trends.

Examination of relationships between the MRI parameters and cognitive performance confirmed that white-matter T1 was shorter (i.e., more “normal”) in subjects who attained higher CFIT scores, whereas gray-matter T1 was unrelated to performance on that test. Unexpectedly, longer gray-matter T1 values were observed in subjects with lower vocabulary scores, although no relationship was found between those scores and the white-matter T1.

The gray–white-matter T1 ratio showed a significant linear relationship with both age and fluid intelligence (Ta-
ence. More than half changes in gray-white-matter regression in the multiple tlo, and hierarchical conducted a TI ratio) were components attributable independent ble I). To partition for fluid (CFIT), the other for crystallized (V3) intelligence. More than half of the variance in CFIT scores \( R^2 \cdot 100\% = 53\% \) was explained by the independent variables \( F(2,23) = 12.94, p < .001 \), with 23% attributable to age alone \( F(1,23) = 11.23, p < .001 \); about 3% was uniquely explained by the gray-white-matter T1 ratio \( F(1,23) = 1.57, \text{n.s.} \), with both variables accounting for 27% of the total variance in CFIT. In contrast, age and the T1 ratio did not explain a significant proportion of variance in V3 scores \( R^2 = .10, F(2,23) = 1.36, \text{n.s.} \). The multiple correlation between CFIT and the set of two independent variables—age and gray-white-matter T1 ratio—was significantly greater than that between V3 and the same pair of predictors: \( Z^* = 2.04, p < .05 \) (Steiger, 1980).

**DISCUSSION**

The results of this study indicate that, at least in terms of molecular properties of tissue water, structural differences between gray and white matter of the medial temporal lobes diminish with age. Indeed, some elderly subjects show very poor differentiation between the two types of cerebral tissue. This finding provides a quantitative confirmation of George et al. ‘s (1981) qualitative observation of reduced gray-white-matter discriminability on CT scans of the elderly. It is also consistent with the reports of decreased gray-white-matter differentiation in the elderly based on rCBF and postmortem studies (Meyer, Kobari, Ichijo, Imai, & Oravez, 1988; Samorajski & Rollsten, 1973). Interestingly, the pattern of age-related decrease in gray-white-matter differentiation is the converse of the diverging pattern observed at the earliest stages of postnatal development (Barkovich, Kjos, Jackson, & Norman, 1988; Holland, Haas, Norman, Brant-Zawadzki, & Newton, 1986), with equality of gray- and white-matter T1 values found only in neonates (Bottomley et al., 1984). Although different processes may be responsible for reduced gray-white differentiation in the elderly, the resemblance of their final product illustrates how the development of the brain may have come full circle in the cycle of growth and decline.

The nonlinear age-related prolongation of the white-matter T1 found in this study is consistent with a similar quadratic trend for the white-matter T2 reported by Jer­nigan et al. (in press). The cubic age trend for hippocampal T1 suggests that whereas the gray-matter T1 in the elderly is shorter than in the young, in the “old” old T1 prolongation is observed. The reason for this trend is unclear. One possibility is that cerebral atrophy resulting in enlargement of spaces filled with cerebrospinal fluid (CSF) may be the dominant process in the older elderly, whereas the brains of their younger peers may be more affected by age-related processes reducing the proportion of free water in the tissue. Indeed, comparison of gray-matter T1 between the young (age < 35) and the very old (age > 68) subjects showed no differences [648.6 msec vs. 629.6 msec; \( t(14) = .79, \text{n.s.} \)], whereas the younger elderly (age 60–68) exhibited a significant shortening of the hippocampal T1 [600.6 msec; \( t(15) = 2.14, p < .05 \)]. Changes in temporal gray and white mat-
ter frequently observed in the aging population may stem from a variety of loosely linked or even completely independent cerebral processes listed in the introduction to this report; a near-zero correlation between $T_1$ times measured in cerebral white and gray matter is consistent with this view. Taken together, however, these measures may summarize the net result of multiple age-related cerebral changes in a single index.

It is unclear whether or not gray-white-matter de-differentiation reflects a general trend in the aging brain and whether or not relationships between aging of specific brain structures and specific cognitive abilities may be established at all. This study provides evidence supporting an association between a general cognitive measure and tissue properties in a selected cerebral region. To assess the validity of this approach and its generality, future studies should include measurements of brain structures and cognitive abilities aimed at double dissociation of structure-function relationships. If the validity of $T_1$ measures as indexes of the extent of age-related cerebral transformations is established, gray-white-matter discriminability may become a promising biological marker of cerebral aging. Provided the relationship between fluid intelligence and such a marker is replicated, we may obtain a powerful method of studying biological foundations of cognitive aging in vivo.

Sprott (1988) listed the following requirements for good biomarkers of aging: (1) be measurable in a noninvasive risk-free procedure, (2) reflect basic physiology and not pathology, (3) show broad generalizability across species, (4) be reproducible within and across laboratories, (5) have a measurable and predictable rate of change, and (6) show significant changes in relatively short periods of time. Evaluation of the proposed index of cerebral aging according to Sprott’s criteria reveals that the gray-white ratio of $T_1$ times completely satisfies only (1), while the rest of these criteria have yet to be met. We believe that careful selection of our sample makes meeting requirement (2) very likely. Requirement (3) is partially satisfied, for gray-white-matter $T_1$ ratio in young laboratory mammals is similar to that measured in humans (Bottomley et al., 1984). We are not aware, however, of animal studies of age-related changes in gray-white-matter differentiation. Such studies are feasible and would be necessary for validation of gray-white-matter $T_1$ ratio as a biomarker of aging. The question of satisfying (4) can be answered only by replication, whereas (5) and (6) require a longitudinal study. If these conditions are met, the gray-white-matter differentiation index will be useful in exploring the biological foundations of cognitive and emotional changes associated with old age, the response of the aging CNS to environmental insults, chronic medication use, and variations in nutrition.

In respect to cognitive correlates of cerebral aging, the results of this study are not clear-cut. A substantial age-related difference in fluid intelligence was observed, as expected, and the zero-order correlation between CFIT scores and the index of gray-white-matter differentiation was moderately high. On the other hand, age alone accounted for a substantial proportion of variance in fluid intelligence, and the unique contribution of the white-gray-matter differentiation index was nonsignificant. It is important to emphasize, however, that the largest share of nonerror variance in fluid intelligence was explained by the commonality between age and the gray and white-matter properties. We could not find a readily interpretable association between $T_1$ values in the hippocampal formation and age-related changes in cognition.

The correlations between the cognitive measures and the $T_1$ values were further examined in a subsample of elderly subjects (age ≥ 60 years). When these subjects were considered separately, the direction and the magnitude of the correlations among age, CFIT, and the white-matter $T_1$ were preserved. The older of the elderly subjects had longer white-matter $T_1$ ($r = .69, p < .01$) and tended to exhibit lower CFIT scores ($r = .52, p < .1$), whereas those with lower CFIT scores tended to have prolonged white-matter $T_1$ ($r = .49, p < .13$). The results of a similar analysis of the correlations among the gray-matter $T_1$, age, and vocabulary scores were quite different. Compared with the pattern observed in the full sample, the one revealed by the analysis of the subsample of the elderly was reversed: Gray-matter $T_1$ tended to become longer, not shorter, with age ($r = .52, p < .1$), and vocabulary scores showed a decreasing, not increasing, age-related trend ($r = -.51, p < .11$); the correlation between $V_3$ and gray-matter $T_1$ dropped to $r = -.20, p > .5$, n.s. Thus, a between-group comparison revealed that although the elderly subjects (≥ 60 years old) had shorter gray-matter $T_1$ than did the younger subjects (≤ 35 years old), they were also blessed with somewhat higher vocabulary scores. Within the elderly group, however, the oldest subjects showed age-related prolongation of the $T_1$ times and a decline in verbal ability. These analyses suggest that the unexpected negative correlation between verbal ability and hippocampal $T_1$ observed in the total sample is, probably, an artifact. The predicted positive relationship between CFIT scores and hippocampal $T_1$ did not materialize.

The reported findings, although preliminary in nature, raise several important questions for future research. Calendar age is not a specific variable; it represents a variety of physiological and cognitive factors that may influence performance on complex tasks, such as fluid intelligence tests. We cannot, therefore, adopt a model postulating that age affects fluid intelligence directly and indirectly via the measured brain variable ($T_1$ ratio). The problem with such a model is that the relationship between age and a single cerebral measure is not unidirectional. Although the calendar age is obviously not influenced by cerebral integrity, the physiological variables that it represents may be. Thus, the appropriate model would be one including age and gray-white $T_1$ ratio as correlated causes of decline in CFIT. Therefore, in spite of the fact that a substantial proportion of variance is explained by age alone, it is the commonality between the measured...
brain variable and those contained under the “age” label that may draw the interest of future investigators. Unlike
an interaction indicating joint effect of the independent
variables on the dependent one, commonality suggests
redundancy of the influence of the independent variable
on the dependent. Further partitioning of the variance hid­
ted in this common part is a challenging task that will
require exploration of additional indexes of neural aging
within a framework of multivariate study. Such indexes
would include electrophysiological and metabolic mea­
sures of brain work, as well as performance on element­
ary information-processing tasks representing fundamen­
tal components of intelligence.

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