Epidemiology and Prevention

Cardiac Index Is Associated With Brain Aging

The Framingham Heart Study

Angela L. Jefferson, PhD; Jayandra J. Himali, MS; Alexa S. Beiser, PhD; Rhoda Au, PhD; Joseph M. Massaro, PhD; Sudha Seshadri, MD; Philimon Gona, PhD; Carol J. Salton, BA; Charles DeCarli, MD; Christopher J. O’Donnell, MD, MPH; Emelia J. Benjamin, MD, ScM; Philip A. Wolf, MD; Warren J. Manning, MD

Background—Cardiac dysfunction is associated with neuroanatomic and neuropsychological changes in aging adults with prevalent cardiovascular disease, theoretically because systemic hypoperfusion disrupts cerebral perfusion, contributing to subclinical brain injury. We hypothesized that cardiac function, as measured by cardiac index, would be associated with preclinical brain magnetic resonance imaging (MRI) and neuropsychological markers of ischemia and Alzheimer disease in the community.

Methods and Results—Brain MRI, cardiac MRI, neuropsychological, and laboratory data were collected on 1504 Framingham Offspring Cohort participants free of clinical stroke, transient ischemic attack, or dementia (age, 61±9 years; 54% women). Neuropsychological and brain MRI variables were related to cardiac MRI-assessed cardiac index (cardiac output/body surface area). In multivariable-adjusted models, cardiac index was positively related to total brain volume (P=0.03) and information processing speed (P=0.02) and inversely related to lateral ventricular volume (P=0.048). When participants with clinically prevalent cardiovascular disease were excluded, the relation between cardiac index and total brain volume remained (P=0.02). Post hoc comparisons revealed that participants in the bottom cardiac index tertile (values <2.54) and middle cardiac index tertile (values between 2.54 and 2.92) had significantly lower brain volumes (P=0.04) than participants in the top cardiac index tertile (values >2.92).

Conclusions—Although observational data cannot establish causality, our findings are consistent with the hypothesis that decreasing cardiac function, even at normal cardiac index levels, is associated with accelerated brain aging. (Circulation. 2010;122:690-697.)

Key Words: brain ■ cardiac output ■ epidemiology ■ imaging ■ neuropsychology ■ atrophy

Cardiac function indexes have been related to neuropsychological impairment1-2 and dementia3 among patients with severe cardiomyopathies. In the absence of end-stage heart disease, less is known about how subclinical cardiac dysfunction affects brain aging. Recent cross-sectional work in referral-based samples of cardiovascular patients free of end-stage heart disease indicates that reduced cardiac output is associated with executive dysfunction4 and increased white matter hyperintensity (WMH).5 These data suggest that even subtle cardiac dysfunction is related to central nervous system injury.

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The mechanism underlying an association between cardiac function and maladaptive brain aging remains unknown, but reduced systemic perfusion may affect cerebral perfusion homeostasis. Convention suggests that autoregulatory processes augment blood flow to the brain during brief periods of reduced cardiac function,6 but data indicate that autoregulatory mechanisms are less effective with subclinical or chronic systemic flow reductions.7,8 Animal data demonstrate that
lowering cardiac output directly reduces cerebral blood flow, and clinical studies indicate that cerebral blood flow increases by >50% after heart transplantation. Such reduced systemic blood flow appears to influence cerebral blood flow homeostasis and may contribute to subclinical brain injury.

Initial research in this area has been restricted to small referral samples with clinical cardiovascular disease (CVD). These studies have not systematically adjusted for environmental risk factors known to contribute to both central nervous system and myocardial injury. The proposed cross-sectional investigation aims to enhance knowledge about relations between cardiac function and brain aging by extending prior work to a large epidemiological cohort, using a more precise cardiac imaging method (MRI) with excellent reproducibility for quantifying cardiac function, and simultaneously considering shared risk factors for central nervous system and myocardial injury. On the basis of prior animal and clinical research, we hypothesized that MRI-assessed cardiac function is associated with cognitive and neuroimaging markers of preclinical Alzheimer disease (ie, decreased learning/memory, brain volume, and hippocampal volume and increased lateral ventricular volume) and cerebrovascular disease (ie, decreased executive functioning/information processing and increased WMH) in a community-based cohort of adults free of clinical dementia or stroke.

Methods

Participants
The Framingham Offspring Study design and selection criteria have been described elsewhere. Briefly, from 1971 to 1975, 5124 participants were recruited and have been examined every 4 to 8 years since. The present sample was derived from 3539 participants who refused or were unable to undergo brain MRI were ancillary neuropsychological and brain MRI study. Those participants who refused or were unable to undergo brain MRI were ancillary neuropsychological and brain MRI study. Those participants who refused or were unable to undergo brain MRI were included in the present analyses (n = 696 of 1504 participants).

Neuropsychological Protocol
As previously described, FHS participants completed a neuropsychological protocol that was based in part on measures administered to the original cohort >25 years ago. In more recent cycle examinations, tests were added to assess cognitive domains not adequately captured by the original protocol. Measures included in the protocol have strong reliability and validity and are sensitive to cognitive functions mediated by both frontal-subcortical and cortical systems. For the present study, neuropsychological performances were adjusted for age and education, separately by sex, and these values were expressed in SD units. Natural logarithmic transformations were applied to normalize raw scores with skewed distributions. Next, each variable was regressed onto age and education categories, and residuals from these regressions were standardized with a z-score transformation. Because the protocol is made up of numerous tests measuring related cognitive domains, a factor analysis was conducted to define composites with the use of oblique rotation. The data reduction process yielded 5 neuropsychological factors, and z-score transformations were applied to each of the 5 factors. The 5 composites and corresponding tests are as follows: (1) verbal memory: Wechsler Memory Scale (WMS) Logical Memory Immediate and Delayed Recall; (2) visuospatial memory: WMS Visuospatial Memory Immediate and Delayed Recall; (3) visual learning: Paired Associate Learning Immediate and Delayed Recall; (4) executive functioning/information processing: Trail Making Test Parts A and B; and (5) language/object recognition: Boston Naming Test, and Wechsler Adult Intelligence Scale-Revised Similarities subtest.

Brain MRI Acquisition
The FHS MRI acquisition protocol has been reported in detail elsewhere. Briefly, the majority of participants were imaged on a Siemens 1-T MR machine (Siemens Medical, Erlangen, Germany) with a T2-weighted double spin-echo coronal imaging sequence. Digital information was postprocessed by a central laboratory blinded to demographic and clinical information. Quantification was performed with a custom-written computer program operating on a UNIX Solaris platform (Sun Microsystems, Santa Clara, Calif). The semiautomated segmentation protocol for quantifying total cranial volume, total brain volume, frontal lobar volume, lateral ventricular volume, hippocampal volume, and WMH has been described elsewhere, as has the interrater reliabilities for these methods. For the present study, intrarater and interrater reliabilities were consistently >0.90. Hippocampal data were available for a subset of the sample included in the present analyses (n = 696 of 1504 participants).

Cardiac MRI Acquisition
The FHS cardiac MRI protocol has been reported in detail elsewhere. Briefly, supine cardiac MRI was performed with a Philips 1.5-T MR scanner (Philips Medical Systems, Cleveland, Ohio) with a 5-element commercial cardiac array coil for radiofrequency signal reception. End-expiratory breath-hold, ECG-gated cine steady-state free-precession images were acquired in 2-chamber, 4-chamber, and contiguous short-axis orientations (temporal resolution, 39 ms; repetition time, R-R interval; echo time, 9 ms; flip angle, 30°; field of view, 400 mm; matrix size, 208 × 256; slice thickness, 10 mm; gap, 0). Steady-state free precession is advantageous for quantifying cardiac function over conventional MRI sequences because it provides briefer acquisition time and enhanced signal-to-noise ratio. Cardiac MRI data analysis was performed by an experienced reviewer blinded to clinical information using dedicated software (EasyVision 5.1, Philips Medical Systems). End-systolic phase was determined as the minimal cross-sectional area of a midventricular...
slice. End-diastolic volume (EDV) and end-systolic volume (ESV) were computed by the summation-of-disks method with cardiac output (in L/min) calculated as follows: (EDV–ESV) × heart rate. Cardiac MRI methods yield more precise measurements of EDV, ESV, and stroke volume compared with echocardiography, and cardiac output measurements are highly reproducible. Intraclass correlation coefficients for EDV and ESV are 0.95 and 0.92, respectively. Interobserver coefficients of variation for EDV and ESV are 2.6% and 3.5%, respectively. Interobserver coefficients of variation for EDV and ESV are 3.5% and 4.8%, respectively. Interobserver coefficients of variation for EDV and ESV are 2.6% and 3.5%, respectively.

Statistical Analysis
To standardize cardiac output values across participants for analytic purposes, cardiac output was divided by body surface area to yield cardiac index measured in liters per minute per meter squared (L/min/m²). Total brain volume, frontal lobe volume, lateral ventricular volume, hippocampal volume, and WMH were expressed as percent of total cranial volume to adjust for head size, yielding percentages for analyses. WMH was natural log–transformed to normalize its skewed distribution. All models included the following covariates, which were selected based on prior work: age, sex, systolic blood pressure, cigarette smoking status, diabetes mellitus, hypertension treatment, atrial fibrillation, and prevalent CVD.

Two primary analyses were conducted. First, multivariable-adjusted linear regressions were used to assess relations between cardiac index and each dependent brain MRI measure (WMH, total brain volume, frontal lobar volume, lateral ventricular volume, hippocampal volume) and the 5 neuropsychological composites (verbal memory, visuospatial memory, verbal learning, information processing/executive functioning, language/object recognition). Post hoc tertile analyses were conducted for significant continuous cardiac index and brain aging relations. Second, to assess how low cardiac index relates to brain aging, multivariable-adjusted linear regression was used to assess relations between low cardiac index (ie, defined as <2.5 L/min/m²; n=456) and brain aging markers.

Secondary analyses were performed excluding participants with prevalent CVD (n=112); using interaction terms to assess the effect modification of relations by sex, age (<60 versus ≥60 years), and ApoE genotype status (ε4 negative versus ε4 positive); and then stratifying analyses by these subgroups. Significance was set at P<0.05 for all models and P<0.10 for analyses assessing effect modification. All data were analyzed with SAS version 9.1 (SAS Institute Inc, Cary, NC).

Results

Participant Characteristics
Participant characteristics are provided in Table 1. The mean sample age was 61 years (34 to 84 years); 54% were women. Low cardiac index (ie, <2.5 L/min/m²) was present in 30% of the sample (n=457). After exclusion of individuals with prevalent CVD (n=112), low cardiac index was still present in 30% of the sample (n=415). Brain MRI mean and SD data and neuropsychological median, minimum, and maximum data are provided in Table 2.

Cardiac Index and Brain MRI
As a continuous measure, cardiac index was positively related to total brain volume (β=0.30, P=0.03) such that an increase in 1 SD of cardiac index resulted in an increase of total brain volume (as a percentage of total cranial volume) of 0.30. As a continuous measure, cardiac index was inversely related to lateral ventricular volume (β=-0.10, P=0.048; ie, as cardiac index increased 1 SD, lateral ventricular volume decreased –0.10). In secondary analyses, when participants with clinically prevalent CVD were excluded from analyses, the significant relation between cardiac index and total brain volume remained (β=0.33, P=0.02); however, the relation between cardiac index and lateral ventricular volume was no longer significant (β=-0.09, P=0.08). Post hoc, cardiac index tertiles were compared to assess changes in brain volume. The top (or highest) tertile of cardiac index was used as the referent for analysis because higher values of cardiac index reflect healthier cardiac function. Participants in the middle cardiac index tertile had an average total brain volume (as a ratio value adjusted for total cranial volume) of 0.35 less than those in the top quartile (β=-0.35, P=0.04). Similarly, participants in the bottom cardiac index tertile had a lower mean total brain volume of 0.36 (β=-0.36, P=0.04) compared with participants in the top tertile, corresponding to an average difference of 1.9 years in brain aging (ie, 0.19 decrease in total brain volume is observed per increased year of age). The Figure depicts the adjusted mean total brain volume with standard error (SE) bars across tertiles of cardiac index. Cardiac index was unrelated to the remaining brain MRI measures when treated as a continuous variable (all P>0.17).

When low cardiac index was related as a dichotomous variable to the brain aging variables, no significant relations were observed between low cardiac index and total brain volume (β=-0.18, P=0.24) or lateral ventricular volume (β=0.02, P=0.68). Cardiac index was unrelated to the remaining brain MRI measures when dichotomized into normal and low subgroups (all P>0.10). See Table 3 for details on all statistical models relating cardiac index to total brain volume.

Table 1. Participant Characteristics
<table>
<thead>
<tr>
<th>Clinical characteristics</th>
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</thead>
<tbody>
<tr>
<td>Age at MRI, y</td>
<td>61±9</td>
</tr>
<tr>
<td>Female, %</td>
<td>54</td>
</tr>
<tr>
<td>Systolic blood pressure, mm Hg</td>
<td>125±18</td>
</tr>
<tr>
<td>Cigarette smoking, %</td>
<td>10</td>
</tr>
<tr>
<td>Diabetes mellitus, %</td>
<td>9</td>
</tr>
<tr>
<td>Atrial fibrillation, %</td>
<td>3</td>
</tr>
<tr>
<td>Hypertension treatment, %</td>
<td>28</td>
</tr>
<tr>
<td>Prevalent CVD, %</td>
<td>7</td>
</tr>
<tr>
<td>Heart failure</td>
<td>6</td>
</tr>
<tr>
<td>Coronary artery disease</td>
<td>2</td>
</tr>
<tr>
<td>Intermittent claudication</td>
<td>1</td>
</tr>
</tbody>
</table>

Table 2. Interval from cycle exam 7 to brain MRI, y | 3.70±1.10 |
| Interval from brain MRI to cardiac MRI, y | 0.68±0.65 |
| Cardiac MRI cardiac index, L/min/m² | 2.77±0.51 |
| Cardiac index tertiles, L/min/m² |       |
| 1 | <2.54  |
| 2 | 2.54–2.92  |
| 3 | >2.92  |
| Impaired cardiac index, % | 30    |

n=1504. Values are given as percentages or mean ± SD as appropriate. CVD does not include clinical stroke or transient ischemic attack. CVD categories are not mutually exclusive.

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Table 2. Brain Aging Descriptive Data

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Mean (SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>WMH</td>
<td>0.09 ± 0.21</td>
</tr>
<tr>
<td>Total brain volume</td>
<td>79.66 ± 3.21</td>
</tr>
<tr>
<td>Frontal lobar volume</td>
<td>36.45 ± 3.33</td>
</tr>
<tr>
<td>Lateral ventricular volume</td>
<td>1.75 ± 1.07</td>
</tr>
<tr>
<td>Hippocampal volume†</td>
<td>0.32 ± 0.05</td>
</tr>
</tbody>
</table>

Neuropsychological data, median (minimum, maximum), total correct unless otherwise noted

F1: verbal memory
- WMS Logical Memory Immediate Recall 12 (2, 21)
- WMS Logical Memory Delayed Recall 11 (0, 21)

F2: visuospatial memory
- WMS Visual Reproduction Immediate Recall 10 (0, 14)
- WMS Visual Reproduction Delayed Recall 9 (0, 14)

F3: verbal learning
- WMS Paired Associate Learning Immediate Recall 14 (3, 21)
- WMS Paired Associate Learning Delayed Recall 9 (3, 10)

F4: information processing/executive function
- Trail Making Test A, time to completion, min 0.5 (0.2, 2.3)†
- Trail Making Test B, time to completion, min 1.2 (0.4, 5.0)‡

F5: language/object recognition
- Hooper Visual Organization Test 26 (3, 30)
- WAIS-R Similarities 17 (2, 25)
- Boston Naming Test, 30-Item 28 (16, 30)

WAIS-R indicates Wechsler Adult Intelligence Scale, Revised. n=1504. WMH is log-transformed.

*All brain MRI variables are adjusted for total cranial volume and presented as percentages.
†Analyses based on n=695.
‡Higher values denote worse performance.

Cardiac Function and Neuropsychological Variables

Cardiac index as a continuous variable was unrelated to any of the neuropsychological factor scores (all \(P>0.23\)). However, low cardiac index (ie, \(<2.5 \text{ L} \cdot \text{min}^{-1} \cdot \text{m}^{-2}\)) was borderline significantly related to the executive function/information processing factor (\(\beta=-0.10, P=0.06\)). Post hoc analyses isolating the individual neuropsychological tests comprising this factor score suggest that low cardiac index was significantly related to poorer performances on the Trail Making Test Part A, assessing information processing speed (\(\beta=-0.13, P=0.02\)). In secondary analyses, when participants with prevalent CVD were excluded, the borderline significant relation between low cardiac index and the information processing/executive function factor was attenuated (\(\beta=-0.11, P=0.05\)).

Effect Modification by Age

There was a significant interaction between age and cardiac index in their relation to total brain volume (\(P=0.01\)) such that the magnitude of the association was stronger in the adults <60 years of age compared with the older adults (Table 3). No additional significant interactions were observed between age and cardiac index in their relation to the brain aging variables.

Effect Modification by Sex

There was a significant interaction between cardiac index and sex in their relation to total brain volume (\(P=0.03\)); the magnitude of the association was stronger for men than women (Table 3). No additional significant interactions were observed between sex and cardiac index in their relation to the brain aging variables.

Effect Modification by ApoE e4 Status

There was a possible interaction between low cardiac index and ApoE status in their relation to the visuospatial memory factor (\(P=0.09\)) such that the magnitude of the association was stronger for e4-positive carriers than e4-negative carriers. No additional significant interactions were observed between ApoE e4 status and cardiac index in their relation to the brain aging variables.

Discussion

In this study of a community-dwelling cohort of ambulatory adults, we found that cardiac index was associated with total brain volume and lateral ventricular volume, both markers of accelerated brain aging\(^{17,42}\); however, when analyses excluded participants with prevalent CVD, only the association between cardiac index and total brain volume remained. The association was stronger for individuals <60 years of age, which coincides with a period in the lifespan with reduced risk for abnormal brain changes, possibly allowing the influence of cardiac index on brain health to be more prominent. Individuals in the top tertile of cardiac index had a higher mean total brain volume equivalent to nearly 2 years of healthy brain aging compared with those participants in either the middle or bottom tertile of cardiac index. Although cardiac index as a continuous variable was unrelated to neuropsychological performances, when clinical cutoffs were applied, low cardiac index was related to information processing speed, a finding that was modestly attenuated when participants with prevalent CVD were excluded from analyses.

Collectively, these results suggest that even in the absence of prevalent CVD, cardiac index is related to total brain volume, a neuroimaging marker of brain health, and low cardiac index is related to information processing speed. Prior research relating cardiac output to neuroimaging and neuropsychological phenotypes of maladaptive brain aging in clinical cohorts has yielded significant associations.\(^2,4,5\) For instance, among patients with severe cardiomyopathies,\(^6\) reduced cardiac output is related to cognitive impairment. Among elders with prevalent CVD, reduced cardiac output is related to both executive dysfunction\(^4\) and WMH.\(^5\)

The present findings enhance prior research in 2 important ways. First, these data extend past reports by demonstrating that cardiac output is related to brain aging markers in the absence of clinical CVD in a community-based cohort. Second, on the basis of the cohort under investigation, the
point at which cardiac index is significantly related to abnormal brain health appears to differ from the clinical threshold for abnormal cardiac function (ie, a cardiac index value 2.5 is often used to define impaired cardiac function40). The Figure illustrates that the level at which cardiac index is associated with differences in brain health (defined as total brain volume) appears to be 2.9, suggesting that a range of normal cardiac index values (ie, 2.5

Figure. Mean total brain volume (with SE bars) adjusted for age, sex, systolic blood pressure, cigarette smoking status, diabetes mellitus, hypertension treatment, atrial fibrillation, and prevalent CVD is depicted by tertile of cardiac index. Tertile 3 is significantly different from tertiles 1 (P=0.04) and 2 (P=0.04).

Table 3. Cardiac Index and Total Brain Volume Results

<table>
<thead>
<tr>
<th></th>
<th>Total Brain Volume</th>
<th>Analyses Stratified by Sex</th>
<th>Analyses Stratified by Age</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Total Sample</td>
<td>Without Prevalent CVD</td>
<td>Female (n=806)</td>
</tr>
<tr>
<td></td>
<td>(n=1504)</td>
<td>(n=1392)</td>
<td></td>
</tr>
<tr>
<td>Heart index</td>
<td>β±SE P</td>
<td>β±SE P</td>
<td>β±SE P</td>
</tr>
<tr>
<td>Continuous</td>
<td>0.30±0.14 0.03</td>
<td>0.33±0.14 0.02</td>
<td>0.07±0.19 0.72</td>
</tr>
<tr>
<td>Tertile 1 (bottom/low)</td>
<td>-0.36±0.17 0.04</td>
<td>-0.41±0.17 0.02</td>
<td>-0.26±0.23 0.25</td>
</tr>
<tr>
<td>Tertile 2</td>
<td>-0.35±0.17 0.04</td>
<td>-0.34±0.17 0.04</td>
<td>-0.32±0.23 0.17</td>
</tr>
<tr>
<td>Tertile 3 (top/high)</td>
<td>Referent</td>
<td>Referent</td>
<td>Referent</td>
</tr>
<tr>
<td>Low cardiac index</td>
<td>-0.18±0.15 0.24</td>
<td>-0.21±0.15 0.17</td>
<td>-0.05±0.19 0.80</td>
</tr>
</tbody>
</table>

β indicates the change in brain volume as a function of a 1-SD change in cardiac index; Cardiac index is cardiac output divided by body surface area, with values <2.5 L·min⁻¹·m⁻² defined as low cardiac index. Low cardiac index, n=457. Models were adjusted for age, sex, systolic blood pressure, smoking status, diabetes mellitus, hypertension treatment, atrial fibrillation, and prevalent CVD. Cardiac index values for tertiles are <2.54 5 L·min⁻¹·m⁻² for tertile 1 (n=501), 2.54 to 2.92 5 L·min⁻¹·m⁻² for tertile 2 (n=502), and >2.92 5 L·min⁻¹·m⁻² for tertile 3 (n=501).
to 2.9) may be related to compromised brain health integrity. These findings require further study, but if replicated, such results may have significant clinical implications, including the early identification of individuals with low (<2.5) or low-normal (2.5 to 2.9) cardiac index for treatment to prevent abnormal brain changes.

The mechanism accounting for associations between cardiac index and markers of brain aging is unknown; however, reduced systemic blood flow may contribute to subclinical brain injury because of its impact on cerebral blood flow homeostasis.\(^4\)\(^3\)\(^4\)\(^5\)\(^6\) Despite autoregulatory mechanisms to preserve blood flow to the brain, research in macaque monkeys has demonstrated that lowering systemic blood flow via reduced cardiac output directly reduces cerebral perfusion.\(^7\) Similar findings have been reported in heart transplant candidates with severe cardiomyopathies; cerebral blood flow values return to healthy levels after restoration of cardiac function.\(^8\) Alterations in cerebral blood flow homeostasis can contribute to clinical or subclinical brain injury by propagating or exacerbating microvascular damage or Alzheimer disease neuropathology. For instance, alterations in cerebral perfusion lead to microvessel structure changes, expression of vascular cell receptors, alterations in microvessel permeability, and vascular remodeling.\(^9\)\(^10\)\(^11\)\(^12\) Furthermore, rats develop Alzheimer disease–related neuropathology after brief cessation of blood flow, including diffuse β amyloid peptide and amyloid precursor protein expression in the hippocampus, entorhinal cortex, and neocortex.\(^13\) In transgenic mouse models of Alzheimer disease, chronic cerebral hypoperfusion places the brain at risk for amyloid deposition, resulting in neuronal death.\(^14\) Thus, reductions in cardiac function and systemic blood flow may lead to subclinical or clinical brain injury by affecting cerebral blood flow homeostasis. Future studies are needed to understand the mechanism(s) accounting for the preliminary epidemiological associations reported here and clinical findings reported elsewhere.\(^2\)\(^4\)\(^5\)\(^6\)\(^7\)

Beyond the primary findings described above, an unanticipated finding was the number of community-dwelling adults with cardiac index values below standard clinical cutoff criteria. That is, ≈30% of participants had low resting cardiac index (ie, values <2.5 L·min\(^{-1}\)·m\(^{-2}\)).\(^4\) When individuals with prevalent CVD were excluded, 30% of participants still had resting cardiac index values <2.5 L·min\(^{-1}\)·m\(^{-2}\). In light of the current observation that cardiac index is associated with cross-sectional markers of accelerated brain aging, such a high proportion of cardiac index values below clinical criteria warrants further investigation.

Our study has a number of strengths, including the large community-based cohort free of clinical dementia and stroke, comprehensive ascertainment of potential confounding variables, an innovative and precise cardiac imaging technique, stringent quality control procedures for measurement of cardiac and brain MRI, and a core reading laboratory for processing measurements. However, the present findings must be tempered by several caveats. First, multiple comparisons were made, raising the possibility of a false-positive finding. Next, the age and racial makeup of the Framingham Offspring Study is predominantly white, of European descent, and middle-aged to elderly, so the generalizability to other races, ethnicities, and age groups is unknown. The exclusion of institutionalized individuals and participants with clinical stroke and the inclusion of individuals willing to undergo MRI yielded a generally healthy sample, thereby reducing the likelihood of finding relations that may be present in the general population that includes individuals with cognitive impairment or stroke. The lack of an association between cardiac index and most of the neuropsychological measures may be due to inadequate power. Similarly, the smaller data set available for analyses relating cardiac index to hippocampal volume may have been insufficiently powered. The observational design limits inferences about causality. The brain MRI data were temporally acquired before the cardiac MRI data, which limits interpretation of directionality. Finally, the cross-sectional design increases the likelihood of residual confounding despite attempts to thoroughly adjust for confounders, including current systolic blood pressure and hypertension medication use, in analytical models. Our results are preliminary, and our findings require replication in other samples.

**Conclusions**

Cardiac index is associated with brain volume, even in individuals without diagnosed prevalent CVD. Although our analyses are based on an observational design and we are unable to establish a causal relation or temporality for the associations observed, we propose that subtle reductions in cardiac index, as well as cardiac index values in the low end of the normal range, may be implicated in accelerating age-related changes in the brain. However, we cannot rule out the possibility that the findings are due to some epiphenomenon. Further investigation into the mechanisms and clinical significance of the association between cardiac index and brain aging is merited.

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**Disclosures**

None.

**References**


**CLINICAL PERSPECTIVE**

This cross-sectional observational study of 1504 community-dwelling, ambulatory adults (age, 34 to 84 years) is among the first epidemiological studies to examine the association between cardiac index and magnetic resonance imaging and neuropsychological markers of brain aging. Individuals in the top tertile of cardiac index (ie, values >2.9 L·min⁻¹·m⁻²) had a significantly higher mean total brain volume, equivalent to nearly 2 years of healthier brain aging, compared with those individuals in the middle (2.5 to 2.9 L·min⁻¹·m⁻²) or bottom (<2.5 L·min⁻¹·m⁻²) cardiac index tertile. These findings suggest that in addition to low cardiac index (<2.5), a range of low-normal cardiac index values (2.5 to 2.9) may be related to compromised brain health integrity (as defined as total brain volume). If a reduced cardiac index is confirmed by other studies to be a risk factor for abnormal brain aging, then early identification of individuals with low or low normal cardiac index values may have important implications for the prevention and treatment of abnormal brain changes, including delaying or preventing the onset of dementia in older adults.
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