Analysis of Genetic Variability and Whole Genome Linkage of Whole-Brain, Subcortical, and Ependymal Hyperintense White Matter Volume

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Background and Purpose—The cerebral volume of T2-hyperintense white matter (HWM) is an important neuroimaging marker of cerebral integrity. Pathophysiology studies identified that subcortical and ependymal HWM are produced by 2 different mechanisms but shared a common risk factor: high arterial pulse pressure. Recent studies have demonstrated high heritability of the whole-brain HMW volume and reported significant and suggestive evidence of genetic linkage. We performed heritability and whole-genome linkage analysis to replicate previous reported findings and to study shared genetic variance, and possible overlap for specific loci, between subcortical and ependymal HWM volumes in a population of healthy Mexican Americans.

Methods—The volumes of subcortical and ependymal HWM regions were measured from high-resolution (1 mm³), 3-dimensional fluid-attenuated inversion recovery images acquired for 459 (283 females, 176 males) active participants in the San Antonio Family Heart Study. Subjects ranged in age from 19 to 85 years of age (47.9±13.5 years) and were part of 49 families (9.4±8.5 individuals per family).

Results—The volumes of whole-brain, subcortical, and ependymal HWM were highly heritable (\(h^2 = 0.72, 0.66, \) and 0.73, respectively). The subcortical and ependymal HWM volumes shared 21% of genetic variability indicating significant pleiotropy. Genomewide linkage analysis showed only a suggestive bivariate linkage for subcortical and ependymal HWM volumes (log of odds = 2.12) on chromosome 1 at 288 cM.

Conclusion—We replicated previous findings of high heritability for the whole-brain HWM volume. We also showed that subcortical and ependymal volume shared a significant portion of genetic variability and the bivariate linkage analysis produced a suggestive linkage near the locus previously identified in a study of whole-brain HWM volume and arterial pulse pressure. (Stroke. 2009;40:3685-3690.)

Key Words: aging ■ brain imaging ■ genetics ■ hyperintense white matter ■ MRI ■ structural imaging ■ white matter disease
lies. It was proposed that genes responsible for mitochondrial functioning, located near this region, may modulate volume of HWM during the normal aging process. The second study performed whole-genome linkage analysis for WB-HWM in a population of 488 hypertensive adults. Univariate linkage analysis only reached suggestive significance and did not overlap with genetic loci reported by Stefano and colleagues, possibly due to its focus on individuals with hypertension. Multivariate analysis reported highly significant loci for WB-HWM and quantitative measurements of hypertension, suggesting a high degree of pleiotropy.

Neither of the previous studies separated the WB-HWM volume into the subcortical and ependymal components. In this article, we pursued 3 goals: (1) to replicate findings of high heritability for WB-HWM volume; (2) to perform a novel analysis of shared genetic variance, and possible overlap for specific loci, between subcortical and ependymal HWM volumes; and (3) to search for chromosomal regions influencing HWM volume in a Mexican American population.

Methods

Subjects and Measurements

Four hundred fifty-nine (283 females, 176 males; average age = 47.9 ± 13.5 years, age range = 19 to 85 years) Mexican American participants in the San Antonio Family Heart Study were recruited for this study. Recruited subjects were from a large extended pedigree composed of 49 families with the average family size of 9.4 ± 8.5 individuals (range = 2 to 38). At the time of the imaging, 144 subjects were treated for hypertension, 80 subjects were treated for Type II diabetes, and 52 subjects had both hypertension and diabetes. Subjects were excluded for MRI contraindications, history of neurological illnesses, or major neurological event (stroke). Subjects’ diagnosis status for hypertension and diabetes were coded as binary covariates. All subjects provided written informed consent on forms approved by the Internal Review Board of the University of Texas Health Science Center at San Antonio.

Imaging was performed at the Research Imaging Center, University of Texas Health Science Center at San Antonio, using a Siemens 3T Trio scanner and an 8-channel head coil. The 3-dimensional T2-weighted imaging data were acquired using a high-resolution (isotropic 1 mm), turbo-spin-echo fluid-attenuated inversion recovery (FLAIR) sequence with the following parameters: TR/TE/TI/flip angle (FLAIR) = 5 seconds/353 ms/1.8 s/180°/221. This 3-dimensional FLAIR protocol was designed to overcome limitation of previous studies of genetics of HWM. The FLAIR sequence was used in prior studies of genetics of HWM. The chosen protocol allowed for sensitive detection of smaller lesions and accurate tracing of lesion boundaries. The 3-dimensional FLAIR sequence applied in prior studies of genetics of HWM. The FLAIR sequence prevents ventricular cerebrospinal fluid pulsation artifacts commonly seen as false-negative hyperintense regions in the 2-dimensional FLAIR sequences.

Measurement of HWM volume from FLAIR images is discussed elsewhere. In short, FLAIR images were preprocessed by removal of nonbrain tissue, registration to the T1-weighted images/Talairach frame, and radiofrequency inhomogeneity correction. HWM regions were manually delineated in 3-dimensional space using in-house software (http://ric.uthscsa.edu/mango) by an experienced neuroanatomist with high ($r^2 > 0.9$) test-retest reproducibility. HWM regions were coded as ependymal regions, contiguous with cerebrospinal fluid structures, and subcortical in accordance with. The WB-HWM volume and the volumes of subcortical and ependymal HWM were measured for each subject.

Genotyping

The details of the genotyping procedure can be found in Kammerer et al. After DNA was extracted from lymphocytes, fluorescently labeled primers from the MapPairs Human Screening set (Versions 6 and 8; Research Genetics, Huntsville, Ala) and polymerase chain reaction were used to amplify 417 microsatellite markers spaced at approximately 10-cM intervals across 22 autosomes. An automated DNA sequencer (ABI Model 377 with Genescan and Genotyper software; Applied Biosystems, Foster City, Calif) was used. The average heterozygosity index for these markers was approximately 0.76. The sex-averaged marker map was confirmed by deCODE genetics and markers not on this map were placed by interpolation based on physical location.

Genotypes were subjected to extensive data cleaning with the SimWalk2 software package. The computation was based on maximum likelihood marker allele frequencies. This statistical procedure is designed to detect inconsistencies and unlikely genotypes. An iterative process was followed to eliminate genotypes that are likely to be erroneous until no more inconsistencies or possible errors remained. After this, the multipoint identity-by-descent matrices were estimated using the Markov chain Monte Carlo methods implemented in Loki. The probabilities of multipoint identity-by-descent allele sharing among all possible pairs of related individuals were computed using the genotypes at all linked markers jointly in the computations.

Heritability and Quantitative Trait Linkage Analysis

Quantitative genetic analyses were performed using a variance components method implemented in SOLAR. SOLAR uses maximum likelihood variance decomposition methods to determine the relative importance of genetic and environmental influences on a trait by modeling the covariance among family members as a function of genetic proximity (kinship). Heritability ($h^2$), the proportion of phenotypic variance accounted for by additive genetic variance ($h^2 = \sigma_g^2/\sigma_p^2$), was assessed by contrasting observed covariance matrices with the covariance matrix predicted by kinship. Bivariate genetic correlation analyses were performed to decompose phenotypic correlations ($r_p$) between regional HWM measurements into the genetic ($r_p g$) and environmental ($r_p e$) correlations for kinship: $r_p = r_p h^2(1/2)(h^2 + 1 - h^2)/1/2 + r_p(1 - h^2)^1/2$. Quantitative trait linkage analysis was performed to localize potential genes influencing phenotypic variation to specific chromosomal locations. Model parameters were estimated using maximum likelihood. The hypothesis of significant linkage was assessed by comparing the likelihood of a classical additive polygenic model with that of a model allowing for both a polygenic component and a variance component due to linkage at a specific chromosomal location. The LOD score given by the log10 of the ratio of the likelihood of the linkage and the polygenic model served as the test statistic for genetic linkage. For this exact pedigree structure and density of markers, an LOD of 1.67 is required for suggestive significance (likely to happen by chance less than once in a genomewide scan) and a LOD of 2.88 is required for genomewide significance at the 0.05 level. Similar to previous studies, HWM volumes were positively skewed. An inverse Gaussian transformation was used to assure normal range for kurtosis and skewness. All genetic analyses were conducted with age, sex, age*sex, age2, age*sex, and diagnostic status for hypertension, diabetes, and heart disorder (encoded as 0 or 1) included as covariates.

Results

Heritability Analysis

WB-HWM volume increased exponentially with age (Figure 1), consistent with previous findings by this center and...
All measures of HWM volume were significantly influenced by genetic factors. Heritability estimate for WB-HWM volume was $0.72 \pm 0.11$ ($P=1.0 \times 10^{-14}$). Consistent with previous reports, the age and age$^2$ covariates were nominally significant ($P<0.10$) for the WB-HWM volume. Lobar, ependymal, and sublobar HWM volumes were also determined to be highly heritable (Table). Age was a significant covariate for both regional HWM volume measurements and age$^2$ approached significance for the ependymal HWM volume. Binary covariates that coded diagnosis for hypertension and diabetes did not reach statistical significance for any of the traits.

**Bivariate Correlation Analysis**

Genetic correlation analysis indicated that the subcortical and ependymal HWM volumes shared 21% of genetic variance ($\rho_G=0.46 \pm 0.12$; $P=0.001$), suggesting some degree of pleiotropy. In contrast, the environmental correlation between HWM measurements was not significant ($\rho_E=-0.07 \pm 0.24$; $P=0.90$). This result suggests that the observed phenotypic correlation between these 2 traits is driven overwhelmingly by shared genes.

**Linkage Analysis**

Genomewide linkage analysis did not reveal a genomewide significant localization for any of the HWM traits. However, suggestive linkage for subcortical and ependymal HWM volumes combined (bivariate) was found (LOD=$2.12$) on chromosome 1 at 288 cM near the p-terminus (Figure 2). Ependymal volume alone also showed suggestive linkage (LOD=$1.72$) at this chromosomal location (Figure 2). WB and subcortical HWM volumes were nominally significant (LOD=$1.51$ and $1.63$, respectively) at this location.

**Discussion**

WB-HWM volume is a complex trait with a large genetic component. Histopathologic findings suggest 2 distinct forms of HWM lesions, subcortical and ependymal. In normal aging, subcortical HWM are thought to result from ischemic and/or neuroinflammatory etiologies. Therefore, formation of

<table>
<thead>
<tr>
<th>Trait</th>
<th>Percent of Total Volume</th>
<th>$h^2$</th>
<th>$p$</th>
<th>Significant Covariates</th>
<th>Variance Explained by Covariates</th>
</tr>
</thead>
<tbody>
<tr>
<td>Whole-Brain</td>
<td>100.00</td>
<td>0.72</td>
<td>1E-14</td>
<td>age (5E-14), age$^2$ (0.08)</td>
<td>28%</td>
</tr>
<tr>
<td>Subcortical</td>
<td>23.57</td>
<td>0.66</td>
<td>4E-11</td>
<td>age (3E-16)</td>
<td>27%</td>
</tr>
<tr>
<td>Ependymal</td>
<td>76.43</td>
<td>0.73</td>
<td>1E-9</td>
<td>age (2E-9), age$^2$ (0.06)</td>
<td>20%</td>
</tr>
</tbody>
</table>
subcortical HWM is thought to be the product of age-related loss of permeability of small vessels, age-related free-radical damage to oligodendrocytes, and immune-system mediated gliosis. In contrast, ependymal HWM appears to be of nonischemic origin. Histopathologic findings indicate that the subependymal HWM is formed by the gliosis of periventricular WM due to the disruptions of the ependymal lining of cerebral ventricles, a condition also commonly observed in traumatic brain injuries. The ependymal gliosis in normal aging is thought to be produced by a condition called pulse-wave encephalopathy, which refers to the mechanical damages caused to the ependymal lining by the pulsatile movements of cerebrospinal fluid due to the intracranial pulse pressure waves that produce “traumatic” microtears in the ependymal lining. The magnitude of the cerebrospinal fluid intracranial pulse pressure waves is linked to the gradient between systolic and diastolic arterial pressure. High arterial pulse pressure was shown to be a major risk factor for vascular damage and small vessel disease and was associated with higher ependymal HWM volumes. Pulse pressure thereby could be a mechanism partially responsible for production of both subcortical and ependymal HWM. In fact, a recent whole-genome study of HWM in hypertensive individuals found evidence of significant multivariate linkage between 2 traits.

Our results in 459 generally healthy Mexican Americans individuals confirmed previous reports of significant genetic control over variation in the WB-HWM volumes. In addition, the heritability estimates for subcortical and ependymal HWM volume measurements were also highly significant. A significant genetic correlation supported the hypothesis that these 2 distinct forms of HWM lesions are partially influenced by common genetic factors. The results of the univariate whole-genome linkage analysis for the WB and regional HWM volumes did not reach statistical significance for linkage. Traits with high heritability estimates do not always produce significant linkages because heritability estimates do not provide information concerning the complexity of the underling genetic architecture. The lack of significant linkage in the presence of significant heritability implies that the WB and regional HWM volumes are polygenic traits with many quantitative trait loci (QTL), each exerting only moderate effects. For example, normal variation in adult height is highly heritable ($h^2$=0.89 to 0.93), but current estimates suggest that up to 44 independent loci are associated with normal stature. In contrast, the heritability of the neuregulin 1 transcript is somewhat lower ($h^2$=approximately 0.50), but linkage analysis indicated a single locus (LOD=15.8) on chromosome 8. However, results from bivariate linkage analysis for subcortical and ependymal volumes achieved suggestive linkage at marker 288 on chromosome 1. This finding suggests that some of the shared genetic variance between 2 traits reported may be related to this chromosomal region. Turner and colleagues reported a suggestive linkage association for bivariate linkage analysis for the WB-HWM volume and pulse pressure (difference between systolic and diastolic blood pressure) market 274 on chromosome 1. This hints on the nature of common genetic variance between 2 pools of HWM. It also supports the previously suggested hypothesis regarding the formation of ependymal gliosis and the link between ependymal and lobar HWM volumes. In contrast, the current study did not replicate findings of significant linkage on chromosome 4 as reported in DeStefano et al. Indeed, at this chromosomal location, our peak LOD score was only approximately 0.3.

It is important to note that lack of overlap in genetic loci among this and other studies may be due to a number of potential issues. Genetic factors vary across different ethnicities. Prior studies of HWM were focused on populations of European ancestry, whereas our study is the first to examine...
Mexican Americans, a population with significant Native American admixture. If relatively rare variants are involved in the determination of quantitative variability, we may expect considerable differences in the localization of the most important genetic loci across populations. Additionally, this study and 2 other studies of HWM are relatively underpowered and are likely to miss important chromosomal localizations. Also, in general, linkage studies of such complex phenotypes cannot be used to exclude genetic regions for important QTLs. Thus, lack of concordance cannot be interpreted as evidence against the hypothesis that a QTL exists in a particular genomic region. Finally, the cross-sectional nature of these data is suboptimal. There are well-known limitations to the conclusions that can be made about longitudinal processes such as aging from cross-sectional measurements. The longitudinal studies often fail to confirm the age-related trends observed in the cross-sectional samples due to heterogeneity of individual aging trends. Longitudinal data from the Australian Stroke Prevention Study reported large intersubject differences in the rates of accumulation of HWM, and subjects diagnosed with neurological disorders were found to have accelerated rates of accumulation of HWM volume. It is unclear to what degree longitudinal study design can confound the genetic analysis of HWM volume. Age and age2 accounted for up to 27% of the variability in this study and others. The individual rates of accumulation of the HWM volume rates greatly vary from subject to subject, possibly due to individual genetic responses to aging (eg, a genotype-by-age interaction). Hence, it may be useful to explicitly allow for the potential influences of genotype-by-age interactions. Although advanced statistical genetic methods for family-based data allow for the formal detection of such interactions within cross-sectional data, longitudinal family studies will have much greater power to localize and ultimately identify the specific genes involved in intersubject differences in accumulation rates of HWM volume.

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Disclosures

None.

References


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