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Exome chip analysis identifies low-frequency and rare variants in *MRPL38* for white matter hyperintensities on brain MRI

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Abstract

Background and Purpose White matter hyperintensities (WMH) on brain magnetic resonance imaging are typical signs of cerebral small vessel disease and may indicate various pre-clinical, age-related neurological disorders such as stroke. Though WMH are highly heritable, known common variants explain a small proportion of the WMH variance. The contribution of low-frequency/rare coding variants to WMH burden has not been explored.

Methods In the discovery sample we recruited 20,719 stroke/dementia-free adults from 13 population-based cohort studies within the Cohorts for Heart and Aging Research in Genomic Epidemiology consortium, among which 17,790 were of European ancestry (EA) and 2,929 of African ancestry (AA). We genotyped these participants at ~250,000 mostly exonic variants with Illumina HumanExome BeadChip arrays. We performed ethnicity-specific linear regression on rank-normalized WMH in each study separately, which were then combined in meta-analyses to test for association with single variants and genes aggregating the effects of putatively functional low-frequency/rare variants. We then sought replication of the top findings in 1,192 adults (EA) with whole exome/genome sequencing data from two independent studies.

Results At 17q25, we confirmed the association of multiple common variants in *TRIM65*, *FBF1*, and *ACOX1* ($p < 6 \times 10^{-7}$). We also identified a novel association with two low-frequency non-synonymous variants in *MRPL38* (lead: rs34136221, $p_{EA} = 4.5 \times 10^{-8}$) partially independent of known common signal ($p_{EA(\text{conditional})} = 1.4 \times 10^{-3}$). We further identified a locus at 2q33 containing common variants in *NBEAL1*, *CARF*, and *WDR12* (lead: rs2351524, $p_{\text{all}} = 1.9 \times 10^{-10}$). Although our novel findings were not replicated due to limited power and possible differences in study design, meta-analysis of the discovery and replication samples yielded stronger association for the two low-frequency *MRPL38* variants ($p_{rs34136221} = 2.8 \times 10^{-8}$).

Conclusions Both common and low-frequency/rare functional variants influence WMH. Larger replication and experimental follow-up are essential to confirm our findings and uncover the biological causal mechanisms of age-related WMH.

Introduction

White matter hyperintensities (WMH) on brain magnetic resonance imaging (MRI) refer to areas of high intensity signal within cerebral white matter on T2-weighted images. These abnormalities are believed to reflect demyelination and axonal loss as a result of chronic ischemia and blood-brain barrier dysfunction caused by cerebral small vessel disease (SVD)¹ and are commonly observed in the aging population.^{2,3} Substantial evidence supports an association between a high WMH burden and an increased risk of stroke, dementia, and death.⁴

The prevalence and severity of WMH increase with advancing age and presence of cardiovascular risk factors, notably hypertension.⁵⁻⁷ Besides, susceptibility to WMH has a large genetic component, with heritability estimates ranging from 55% to 80%.⁸⁻¹⁰ Genome-wide association studies (GWASs) have identified common genetic variants on chromosome 17q25 associated with WMH.^{11,12} These findings were consistently confirmed in independent studies.¹³⁻¹⁶ Other loci were also identified genome-wide significant in a multi-ethnic GWAS, including common variants at 10q24, 2p21, 1q22, and 2p16.¹² Though additional associations are yet to be discovered, common variants are estimated to account for at most a quarter of the WMH phenotypic variance.¹⁷ The remaining heritability is still unexplained.

Putatively functional, low-frequency (minor allele frequency (MAF) 1~5%) and rare variants (MAF≤1%) within the protein-coding region of the genome (exome), which are not well captured by GWAS arrays, have been proposed to play an important role in complex traits.^{18,19} Yet their effect on WMH has not been explored. Taking advantage of an international collaboration of 13 population-based cohort studies within the Cohorts for Heart and Aging Research in Genomic Epidemiology (CHARGE) consortium,²⁰ we conducted exome-wide association analysis in over 20,000 participants of European or African ancestry genotyped with

Illumina HumanExome BeadChip (exome chip) to identify novel coding variants influencing WMH burden.

Methods

Summary data for this meta-analysis will be available through the database of Genotypes and Phenotypes (dbGaP) CHARGE Summary Results site,²¹ which can be downloaded via authorized access.

Study sample

Thirteen cohort studies within CHARGE consortium were included in the discovery sample (Supplemental Methods). They followed standardized procedure for subject inclusion, genotype calling, phenotype harmonization, covariate selection, and study-level analysis. In addition, two samples of European ancestry from the Three-City Dijon (3C-Dijon) Study and the Alzheimer's Disease Neuroimaging Initiative (ADNI) were used for replication (Supplemental Methods). Study participants were included in the analyses if they had phenotype, genotype and covariate data available and did not have stroke/dementia at the time of MRI scan. Institutional review boards approved all participating studies, and study participants provided written informed consent.

MRI scan and WMH measurement

MRI scans were performed in each study separately, following a standard procedure (Supplemental Methods). In brief, the magnetic field strength of the scanners used in different studies ranged from 1.5 to 3.0 tesla, except for a single site in the Cardiovascular Health Study (CHS) where the strength was 0.35 tesla. T2-weighted spin-echo pulse sequence were used and complemented by either proton density or fluid-attenuated inversion recovery sequence to contrast WMH against cerebrospinal fluid signal. Axial images were acquired and WMH were

estimated on a semi-quantitative visual rating scale (Atherosclerosis Risk in Communities (ARIC) Study Visit 3 (V3) and CHS) or using a quantitative volumetric method (other studies). The two methods have been compared within ARIC and CHS, and showed high agreement with each other.^{22,23}

Genotyping and quality control

Exome chip is a genotyping array focusing on ~250,000 mostly coding variants discovered through exome sequencing in ~12,000 individuals and observed at least three times across at least two existing sequence datasets, including non-synonymous, splicing, stop-altering variants, most of which are rare (http://genome.sph.umich.edu/wiki/Exome_Chip_Design). Samples from all discovery cohorts were genotyped with Illumina HumanExome BeadChip. Variant calling and quality control in these studies were performed either jointly²⁴ or individually following the same protocol (Supplemental Methods). For the replication samples, whole exome sequencing (WES) was performed only on 3C-Dijon individuals with the extremes of the WMH distribution, according to its study design; Whole genome sequencing was performed on ADNI subjects (Supplemental Methods).

Statistical analysis

Figure 1 shows a flow chart illustrating the analytical approach implemented in this study. Detailed description of the analytical methods can be found in Supplemental Methods. In brief, at the individual study level, we performed ethnicity-specific linear regression on rank-normalized WMH in each study separately, which were then combined in meta-analyses to test for association with single variants and genes aggregating the effects of putatively functional low-frequency/rare variants. Analyses conditioning on hypertension status and lead single nucleotide polymorphisms (SNPs) in previous GWAS were also performed, respectively

(Supplemental Methods). The statistical method utilized in the replication analysis for exome-wide significant variants identified is described in Supplemental Methods.

Annotation and functional interpretation

The predicted consequences of all single variants were obtained via dbNSFP,²⁵ which were then used to aggregate putatively functional variants in the gene-based analysis. Functional prediction scores for the exome-wide significant variants from SIFT,²⁶ PolyPhen,²⁷ and CADD²⁸ were obtained via Ensembl Variant Effect Predictor.²⁹ We also used the Genotype-Tissue Expression (GTEx) database to investigate whether these variants affect gene expression in multiple brain tissues.³⁰

Results

After exclusion and quality control, the discovery sample included 17,790 participants of European ancestry (EA, mean age = 66 years) and 2,929 of African ancestry (AA, mean age = 62 years). Descriptive statistics of the participants from each discovery cohort are summarized in Supplemental Table I. A summary of variants included in the meta-analysis by chromosome is shown in Supplemental Table II.

Our primary analysis identified several common and low-frequency/rare variants associated with WMH ($p < 6 \times 10^{-7}$) in the EA and combined samples but not in the AA sample. They include six variants at 17q25 (four common variants are known and two low-frequency variants in *MRPL38* at 17q25 are novel) and three common variants at 2q33 (Table 1, Figure 2, Supplemental Figure I). No other GWAS loci were significant in our analysis (Supplemental Results). Gene-based analysis showed that *MRPL38* was associated with WMH ($p < 3.6 \times 10^{-6}$) in the EA and combined samples but not in the AA sample (Supplemental Table III, Supplemental Figure II). The association was consistent across studies (Supplemental Table IV). However,

among all variants aggregated in this gene, the two significant variants in the single variant analysis contributed most (Supplemental Table V).

Adjustment for hypertension status did not meaningfully affect the signals in both single variant and gene-based analysis (Table 1, Supplemental Table III). When adjusting for rs7214628, the lead known GWAS SNP, 17q25 associations were no longer significant ($p > 0.05$), except those in *MRPL38*. The two low-frequency variants in *MRPL38* remained nominally significant, with p-value ~ 0.001 in the EA sample and 0.006 in the combined sample (Supplemental Table VI). The same trend was observed in the gene-based analysis (Supplemental Table III).

The replication samples included WES data from 498 individuals with extremes on WMH scale and WGS data from 694 individuals (Supplemental Table VII). We tested nine exome-wide significant variants identified in the discovery sample for replication, which represented three independent signals. Thus the significance threshold was set to $p < 0.017$. The results are shown in Supplemental Table VIII. Four known variants in *TRIM65*, *FBF1*, and *ACOX1* at 17q25 were significant in the 3C-Dijon sample. Two low-frequency variants in *MRPL38* at 17q25 and three common variants at 2q33 were not significant. However, the direction of the association for these variants was the same between the discovery and replication samples, and meta-analysis of the p-values for the discovery and replication samples yielded more significant results for the two variants in *MRPL38*. Test of heterogeneity between discovery and replication samples yielded negative results.

Functional predictions for the nine exome-wide significant variants by SIFT, PolyPhen, and CADD are summarized in Supplemental Table IX. Of note, most 17q25 variants have a scaled CADD score > 10 , indicating they are predicted to be among the 10% most deleterious variants in

the genome. Interestingly, the two low-frequency *MRPL38* variants have the highest scaled CADD score among our top variants (23.9 and 32 for rs34136221 and rs9191, respectively), meaning that they are among the 1% and 0.1% most deleterious variants, respectively. Moreover, both variants are predicted deleterious by SIFT, and rs9191 is predicted probably damaging by PolyPhen.

The expression quantitative trait locus (eQTL) results for these top variants from GTEx database are summarized in Supplemental Table X. In brief, common variants at 17q25 are significant eQTLs for their nearby genes (*FBF1*, *MRPL38*, *TRIM47*, and *TRIM65*) in multiple brain tissues. One of the two low-frequency variants in *MRPL38*, rs34136221, acts as a cis-eQTL for *TRIM47* in cerebellum ($p=1.8\times 10^{-6}$) (Supplemental Figure III (A)). At 2q33, two common variants, rs72932557 and rs35212307, are associated with the expression level of *ICA1L* in frontal cortex ($p=2.6\times 10^{-5}$) (Supplemental Figure III (B)).

Discussion

In this meta-analysis of association studies between WMH burden and exome chip genotypes in 13 community-based cohorts of stroke/dementia-free adults of European and African ancestry, we identified both common and low-frequency/rare variants significantly associated with WMH. At the known 17q25 locus, four exome-wide significant common variants were identified in our previous GWAS because they were true signals and the majority of the participants in our exome chip analysis were also included in the previous GWAS. Meanwhile, we showed that the association between the low-frequency non-synonymous variants in *MRPL38* and WMH was partially independent of the known GWAS signal. Although this gene was reported previously for WMH,^{12,16} the variants identified were mostly common and within the linkage disequilibrium

(LD) block of the lead GWAS SNP at 17q25. According to the design of exome chip, which focuses on different genomic regions and allele frequency spectrum from those for GWAS, it is unlikely that the overlapping sample used in the current study and previous GWAS contributed to the novel low-frequency variants identified here. We also identified significant association at 2q33. In our previous GWAS, this locus (rs72934505, $r^2 > 0.96$) was only suggestively associated with WMH,¹² though it reached genome-wide significance when combining evidence from an additional GWAS of WMH in stroke patients.³¹ We failed to replicate the novel low-frequency variants in *MRPL38* at 17q25 or in those genes at 2q33, possibly due to the low frequency of the variants (*MRPL38*), the small effect size (2q33), the limited sample size, and the difference in sample selection strategy, phenotypic transformation, or regression model used in the replication samples. Using the 3C-Dijon data, we calculated that we had less than 15% power to detect association for the two low-frequency variants in *MRPL38*, less than 7% for the three variants at 2q33, comparing to over 70% power to detect association for those known variants at 17q25 (*TRIM65*, *FBF1*, and *ACOX1*). However, we confirmed the direction of their effect, and indeed, the improved p-values by meta-analyzing the discovery and replication samples as well as no evidence of heterogeneity provided additional support for the novel association of *MRPL38* variants with WMH (Supplemental Table VIII).

The two low-frequency variants in *MRPL38*, rs34136221 and rs9191, are in complete LD with each other. Both have MAF of ~1.6% in our sample, which are consistent with allele frequencies observed in the 1000 Genomes Project. In this study, these low-frequency variants contributed primarily to the significance of the gene-based association of *MRPL38*, which encodes the mitochondrial 39S ribosomal protein L38 containing 380 amino acids. Both variants are non-synonymous. The “A” allele of rs34136221 leads to a change from arginine to

tryptophan at position 99 (R99W), and the “G” allele of rs9191 changes aspartic acid at position 371 to histidine (D371H). These changes are predicted to be deleterious to the protein function by SIFT and CADD (Supplemental Table IX). This protein is an important constituent of the large subunit of mitochondrial ribosomes responsible for assembling mitochondrial DNA-coded proteins essential for oxidative phosphorylation.³² The L38 protein maintains the core architecture of the large subunit’s central protuberance, which interacts with the small subunit and with mitochondrial transfer RNAs bound to the ribosome, and is hence critical for mitochondrial translation.³² The exact function of *MRPL38* is still not clear, but interestingly, the structure of the L38 protein is similar to the phosphatidylethanolamine-binding proteins (PEBPs). PEBPs are identified in numerous tissues and have various functions, including a role in neural development and differentiation, which have been implicated in Alzheimer’s disease and gliomas.³³ Evidence is also accumulating that mutated mitochondrial ribosomal protein genes are involved in impaired mitochondrial translation leading to several neurological diseases.³⁴ For instance, *MRPL18*, another mitochondrial ribosomal protein gene that was found to be upregulated in active multiple sclerosis lesions,³⁵ was also differentially expressed in brains of young spontaneously hypertensive stroke-prone rats and associated with WMH.¹⁶ In addition to the deleterious effect on its encoding protein, rs34136221 also acted as a cis-eQTL for *TRIM47* in cerebellum (Supplemental Figure III (A)), a gene in which common variants have been associated with WMH in previous GWASs,^{11,12} indicating that this variant may mediate the common effect of *TRIM47* on WMH. Taken together, these data provide biological plausibility supporting the role of deleterious non-synonymous variants in *MRPL38* in the pathophysiologic changes in the white matter.

Three significant common variants at 2q33 are in high LD with each other ($r^2 > 0.98$) with a MAF of ~12% in our sample. rs2351524 is located at the 5' untranslated region of *NBEAL1*. This variant was associated with coronary artery disease³⁶ and is likely to affect transcription factor binding.³⁷ *NBEAL1* encodes the neurobeachin-like 1 protein, one of nine Beige and Chédiak-Higashi domain-containing proteins (BDCPs).³⁸ The functions of BDCPs remain largely unknown, but BDCP gene mutations may affect lysosome size, apoptosis, autophagy, granule size, or synapse formation.³⁹ Though NBEAL1 is the least studied and understood BDCP, its coding gene is highly expressed in brain and upregulated in glioma.³⁸ The association of an *NBEAL1* variant reported here is in line with our previous observation that many loci significantly associated with WMH contain variants in genes implicated in malignant brain tumors of the white matter that involve glial cells and further suggests that the genetic pathway may be shared between WMH in aging and glioma, perhaps, through glial cell activation or oxidative DNA damage.⁴⁰ rs72932557 is a non-synonymous SNP in *CARF*. This gene encodes a calcium responsive transcription factor, a DNA-binding protein that modulates the transcription of brain-derived neurotrophic factor (BDNF).⁴¹ BDNF is a nerve growth factor that promotes neuronal survival in the adult brain, and a BDNF polymorphism has been associated with WMH in healthy elderly population.⁴² Animal studies have shown that knockdown of Carf reduced cortical BDNF expression and impaired memory.⁴³ Whether the observed association of the *CARF* variant with WMH is direct or through its effect on BDNF remains to be tested. The recent finding that astrocyte-derived BDNF may promote oligodendrogenesis and recovery from white matter damage is consistent with such a hypothesis.⁴⁴ rs35212307 is a non-synonymous SNP in *WDR12*. This gene encodes the WD repeat domain 12, a component of the PeBoW complex required for ribosome biogenesis.⁴⁵ Variants in *WDR12* have been identified in GWAS

of early myocardial infarction⁴⁶ and coronary artery disease⁴⁷ and have been associated with carotid intima-media thickness.⁴⁸ The relationship between carotid atherosclerosis and WMH has long been reported.⁴⁹ In a recent meta-analysis of 10 population-based studies, carotid atherosclerosis was strongly associated with the presence of WMH (odds ratio =1.42, $p<0.0001$),⁵⁰ suggesting a possible role of *WDR12* on WMH through atherosclerosis. However, this hypothesis was not supported by a recent large meta-analysis of more than 50,000 participants with carotid intima-media thickness, in which exome chip variants in *WDR12* were not significant.⁵¹ In the eQTL analysis by GTEx, both rs72932557 and rs35212307 were associated with the expression level of *ICA1L* in frontal cortex (Supplemental Figure III (B)), a gene suggestively associated with neuroticism.⁵²

Strengths of our study include the large community-based sample of middle-aged to older adults free from stroke/dementia and the first investigation of low-frequency/rare putatively functional variants in the coding regions of the genome in relation to WMH. Several limitations should be noted. In particular, the use of a rank-based inverse normal transformation of WMH did not allow for meaningful estimates of effect size, while having the advantages of harmonizing measures of WMH taken on different scale among individual studies, as well as maintaining desirable statistical properties of the analytical models for low-frequency/rare variant analysis. It should also be noted that, despite close agreement between visual rating and volumetric measurement of WMH, there are still known discrepancies in certain circumstances.⁵³ Possible inconsistent definitions of dementia used to filter out participants across studies as well as presence of correlated SVD traits such as lacunes and microbleeds may represent sources of heterogeneity in our sample, which may have diminished our power to detect genetic effects. In addition, variants captured by exome chip represent only a fraction of the genetic diversity of the

genome; we were unable to examine even rarer variants or those that lie outside the coding regions. Whole genome sequencing can provide a more comprehensive view of the genome. Recently, an initial WGS effort for brain imaging traits was made in the Framingham Heart Study, in which the investigators identified an independent common signal at 17q25 (rs9889965) associated with WMH volume.⁵⁴ Intriguingly, consistent with our novel finding of low-frequency *MRPL38* variant (rs34136221), this variant is also an eQTL for *TRIM47* in brain tissues, which provides additional evidence for the role of eQTLs in the development of WMH through the expression of *TRIM47* involved in glial cell activation. The WGS study also identified two novel loci. While confirmation of these loci in additional samples is needed, they were not observed in our data (not shown).

In conclusion, in this exome chip analysis of over 20,000 stroke/dementia-free individuals of European or African ancestry, we identified low-frequency non-synonymous variants in *MRPL38* at 17q25 associated with WMH partially independently of known common signals. In addition, we showed that common variants at 2q33, a previous suggestive locus in stroke-free populations, reached exome-wide significance in our analysis. Functional annotation provides further support for a role of pathways involved in glial cell activation and oxidative damage, possibly common to WMH and glioma. Larger replication and experimental follow-up are essential to confirm our findings and uncover the biological causal mechanisms of age-related WMH, and thus to provide a foundation for identifying targets for diagnosis, treatment and prevention.

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Figure Legends

Figure 1. A flow chart illustrating the analytical approach implemented in this study. MRI: magnetic resonance imaging; WMH: white matter hyperintensities; ICV: intracranial volume; HT: hypertension; SNP: single nucleotide polymorphism; PC: principal component; GWAS: genome-wide association study; EA: European ancestry; AA: African ancestry. *Field center was adjusted if available; Total ICV was adjusted in cohorts with volumetric measure of WMH; Family relationship was adjusted in cohorts with family data; HT was adjusted in the conditional analysis only. †GWAS lead SNP was adjusted in the conditional analysis assessing independence of significant findings at known loci only.

Figure 2. Manhattan plots for single variant analysis without adjustment for hypertension status in the sample of (A) European ancestry (EA), (B) African ancestry (AA), and (C) their combination (EA + AA), respectively. The minor allele frequency threshold was >0.1%. The significance threshold was $p < 6 \times 10^{-7}$ (grey horizontal line). Two significant loci (17q25 and 2q33) in the EA and combined sample were highlighted in green.

Tables

Table 1. Summary of exome-wide significant variants in either EA, AA, or combined sample.

rsID	Locus	chr:pos	Gene	Function	RA	Sample	N	RAF	p (Unadjusted)	p (HT-adjusted)
rs3760128	17q25	17:73886888	<i>TRIM65</i>	NS	G	EA	17,790	0.324	+8.18E-10	+3.50E-10
						AA	2,747	0.711	+7.06E-02	+7.65E-02
						All	20,537	0.375	+1.54E-10	+7.14E-11
rs2351524	2q33	2:203880992	<i>NBEAL1</i>	UTR5	T	EA	17,103	0.123	-5.67E-09	-1.83E-08
						AA	2,323	0.062	-4.25E-03	-5.28E-03
						All	19,426	0.115	-1.92E-10	-7.43E-10
rs2305913	17q25	17:73922941	<i>FBF1</i>	NS	C	EA	17,790	0.327	+6.47E-09	+3.42E-09
						AA	2,747	0.679	+3.92E-01	+4.19E-01
						All	20,537	0.374	+8.27E-09	+5.03E-09
rs1135640	17q25	17:73949540	<i>ACOX1</i>	NS	G	EA	17,422	0.332	+2.21E-08	+1.03E-08
						AA	2,929	0.687	+3.97E-01	+4.32E-01
						All	20,351	0.383	+2.83E-08	+1.57E-08
rs72932557	2q33	2:203846817	<i>CARF</i>	NS	T	EA	17,790	0.123	-3.49E-08	-6.53E-08
						AA	2,929	0.034	-1.32E-01	-2.65E-01
						All	20,719	0.11	-1.13E-08	-3.44E-08
rs35212307	2q33	2:203765756	<i>WDR12</i>	NS	C	EA	17,790	0.123	-3.85E-08	-1.11E-07
						AA	2,929	0.033	-1.05E-01	-1.46E-01
						All	20,719	0.11	-1.09E-08	-3.89E-08
rs34136221	17q25	17:73898188	<i>MRPL38</i>	NS	A	EA	17,790	0.016	+4.47E-08	+4.15E-08
						AA	2,929	0.004	-7.69E-02	-1.06E-01
						All	20,719	0.014	+6.11E-07	+4.93E-07
rs9191	17q25	17:73894963	<i>MRPL38</i>	NS	G	EA	17,790	0.016	+1.76E-07	+1.71E-07
						AA	2,929	0.004	-1.25E-01	-1.71E-01
						All	20,719	0.014	+1.62E-06	+1.36E-06
rs1135889	17q25	17:73926121	<i>FBF1</i>	NS	A	EA	17,790	0.221	+4.72E-07	+2.08E-07
						AA	2,929	0.194	-7.73E-01	-8.32E-01
						All	20,719	0.217	+4.03E-06	+1.75E-06

EA: European ancestry; AA: African ancestry; RA: risk allele; RAF: risk allele frequency; HT: hypertension; NS: non-synonymous; UTR5: 5' untranslated region. The sign of the p-values indicates the direction of the risk allele.