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Genetic Associations With White Matter Hyperintensities Confer Risk of Lacunar Stroke

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- *Background and Purpose*—White matter hyperintensities (WMH) are increased in patients with lacunar stroke. Whether this is because of shared pathogenesis remains unknown. Using genetic data, we evaluated whether WMH-associated genetic susceptibility factors confer risk of lacunar stroke, and therefore whether they share pathogenesis.
- *Methods*—We used a genetic risk score approach to test whether single nucleotide polymorphisms associated with WMH in community populations were associated with magnetic resonance imaging–confirmed lacunar stroke (n=1,373), as well as cardioembolic (n=1,331) and large vessel (n=1,472) Trial of Org 10172 in Acute Stroke Treatment subtypes, against 9,053 controls. Second, we separated lacunar strokes into those with WMH (n=568) and those without (n=787) and tested for association with the risk score in these 2 groups. In addition, we evaluated whether WMH-associated single nucleotide polymorphisms are associated with lacunar stroke, or in the 2 groups.
- **Results**—The WMH genetic risk score was associated with lacunar stroke (odds ratio [OR; 95% confidence interval [CI]]=1.14 [1.06–1.22]; P=0.0003), in patients both with and without WMH (WMH: OR [95% CI]=1.15 [1.05–1.26]; P=0.003 and no WMH: OR [95% CI]=1.11 [1.02–1.21]; P=0.019). Conversely, the risk score was not associated with cardioembolic stroke (OR [95% CI]=1.03 [0.97–1.09]; P=0.63) or large vessel stroke (OR [95% CI]=0.99 [0.93,1.04]; P=0.39). However, none of the WMH-associated single nucleotide polymorphisms passed Bonferroni-corrected significance for association with lacunar stroke.
- *Conclusions*—Genetic variants that influence WMH are associated with an increased risk of lacunar stroke but not cardioembolic or large vessel stroke. Some genetic susceptibility factors seem to be shared across different radiological manifestations of small vessel disease. (*Stroke*. 2016;47:1174-1179. DOI: 10.1161/STROKEAHA.115.011625.)

Key Words: cerebral small vessel diseases ■ genetics ■ genetic association studies ■ leukoencephalopathies ■ stroke, lacunar

Cerebral small vessel disease (SVD) affects the small perforating arteries of the brain and is characterized radiologically by several features, including white matter hyperintensities (WMH), subcortical lacunar infarcts, intracerebral hemorrhages, and cerebral microbleeds.¹ Despite the considerable impact of SVD on health through increased risk of stroke and vascular dementia, the pathophysiological mechanisms underlying SVD remain largely unknown. Pathological findings in diseased vessels include lipohyalinosis and microatheroma,^{2,3} whereas in the parenchyma findings include myelin pallor, enlargement of perivascular spaces, and gliosis.⁴ Many of these findings are common

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to both WMH and lacunar stroke.⁵ However, pathological studies have been hampered by methodological and phenotypic inconsistencies.⁵ In addition, little is known about the extent to which underlying pathogenesis is shared across the radiological manifestations. WMH are increased in lacunar stroke^{6,7}; more so than in other pathological subtypes of stroke, which may indicate that shared pathological processes underlie the 2. In addition, both confluent WMH and lacunar infarcts are a common finding in Mendelian forms of SVD such as cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy,8 although the underlying arterial pathology is considerably different to that of sporadic SVD. However, aside from these exceptions and shared cardiovascular risk factors such as hypertension, few molecular processes have been robustly shown to impact on both lacunar stroke and WMH.

Genetic studies can provide novel insights into SVD and the nature of the relationship between its manifestations. In particular, genome-wide association studies have recently identified multiple genetic variants associated with WMH in community-dwelling individuals⁹ and have been used to show that common variants in *COL4A2*, a gene associated with monogenic SVD, influence sporadic SVD.¹⁰ In addition, genome-wide association studies provide a means of interrogating the relationship between complex traits and assessing whether such traits share pathogenesis. Polygenic risk score approaches can be used to investigate whether 2 conditions are genetically related by testing whether the cumulative effect of trait-associated single nucleotide polymorphisms (SNPs) associated with the first trait influence a second trait. Such approaches have previously been used to assess the influence of risk factors on stroke^{11,12} and to assess the shared susceptibility between stroke and migraine.¹³

In this analysis, we evaluated the impact of common genetic variants associated with WMH from community populations on the risk of lacunar stroke in a well-characterized population of magnetic resonance imaging (MRI)-confirmed lacunar stroke cases and controls. As heterogeneity in the pathology underlying lacunar stroke has been hypothesized,¹⁴ and to test whether an association with lacunar stroke was present in individuals without substantial WMH, we separated our lacunar stroke cases into those with substantial WMH and those with no or mild WMH, testing the influence of WMH-associated variants on these subgroups, as well as on cardioembolic and large vessel strokes. We first used a genetic risk score approach to evaluate the overall evidence that WMH-associated variants affect stroke phenotypes, and then second evaluated whether each of the specific genetic variants is associated with lacunar stroke in both the groups with and without WMH.

Materials and Methods

Study Participants

The study data set consisted of stroke cases obtained from hospital admissions in the UK and Germany (DNA-lacunar, Genes and Ischaemic Stroke [GENESIS] and Wellcome Trust Case Control Consortium 2 [WTCCC2] study), Australia (Australian Stroke Genetics Collaborative [ASGC]), Italy (Milano—Besta Stroke Register [BSR]), and Belgium (Leuven Stroke Study [LSS]), as well as 9053 controls consisting of ancestry-matched individuals from each of the respective case populations (Table). These data sets have been described in detail in previous publications.^{15–18} Genotyping and

 Table.
 Association of SNPs Associated With WMH in Lacunar Stroke Phenotypes

					OR (95% Cl); <i>P</i> Value		
SNP	Gene Region	CHR	BP (hg19)	RA	Lacunar Stroke	Lacunar Stroke w/ WMH	Lacunar Stroke w/out WMH
rs7214628	TRIM65	17	73882148	G	1.02 (0.90–1.15); 0.78	1.08 (0.92–1.28); 0.33	0.94 (0.80–1.11); 0.47
rs72848980	NEURL	10	105319409	G	1.13 (0.99–1.29); 0.08	1.15 (0.97–1.38); 0.12	1.10 (0.92–1.30); 0.30
rs7894407	PDCD11	10	105176179	Т	1.04 (0.94–1.15); 0.49	1.03 (0.90–1.18); 0.65	1.01 (0.88–1.15); 0.89
rs12357919	SH3PXD2A	10	105438112	Т	1.13 (0.99–1.29); 0.07	1.13 (0.95–1.35); 0.15	1.12 (0.94–1.33); 0.19
rs7909791	SH3PXD2A	10	105613178	Α	1.14 (1.03–1.27); 0.01*	1.15 (1.01–1.32); 0.04*	1.19 (1.04–1.36); 0.01*
rs78857879	EFEMP1	2	56135099	Α	0.98 (0.82–1.15); 0.77	1.03 (0.83–1.28); 0.77	0.90 (0.72–1.13); 0.37
rs2984613	PMF1-BGLAP	1	156197380	С	1.10 (0.99–1.22); 0.07	1.13 (0.99–1.30); 0.07	1.06 (0.93–1.20); 0.41
rs11679640	HAAO	2	43141485	С	0.91 (0.81–1.03); 0.15	0.89 (0.76–1.04); 0.13	0.95 (0.81–1.12); 0.55
rs72934505	NBEAL1	2	203916487	Т	1.24 (1.07–1.45); 0.004*	1.29 (1.05–1.59); 0.01*	1.19 (0.98–1.45); 0.08
rs17148926	LOC10050584	5	121510586	С	0.87 (0.76–1.00); 0.05	0.87 (0.73–1.04); 0.14	0.88 (0.74–1.05); 0.17
rs941898	EVL	14	100599437	G	1.12 (1.01–1.25); 0.04*	1.09 (0.95–1.26); 0.22	1.14 (0.99–1.30); 0.07
rs6942756	AHCYL2	7	128886821	G	1.07 (0.96–1.20); 0.22	1.05 (0.91–1.22); 0.50	1.11 (0.96–1.28); 0.15
rs2883428	XM_0039600	1	239571364	G	1.00 (0.89–1.11); 0.96	1.04 (0.90–1.20); 0.62	0.98 (0.85–1.14); 0.83
rs962888	C1QL1	17	43059071	G	1.04 (0.94–1.17); 0.44	1.09 (0.94–1.26); 0.24	1.00 (0.87–1.15); 0.99
rs9515201	COL4A2	13	111040798	Α	1.15 (1.03–1.27); 0.01*	1.13 (0.99–1.30); 0.07	1.14 (1.00–1.31); 0.06

Note: odds ratio are oriented to reflect direction of association with WMH (ie, the reference allele is the risk allele of WMH). BP indicates base position; CHR, chromosomes; CI, confidence interval; hg19, human genome reference 19; OR, odds ratio; RA, reference allele; SNP, single nucleotide polymorphism; and WMH, white matter hyperintensities. **P*<0.05 and in same direction of effect as WMH genome-wide association studies.

imputation of the individuals are described in the online-only Data Supplement. Briefly, all data sets were genotyped on commercially available Illumina arrays and imputed to 1000 Genomes phase 3 using SHAPEIT v2 (for phasing) and IMPUTE v2.2.2 (for imputation).

Phenotype Classification

Subtyping of the ASGC, WTCCC2, GENESIS, BSR, and LSS groups was initially performed using the Trial of Org 10172 in Acute Stroke Treatment (TOAST) classification,19 using clinical assessment as well as brain and vascular imaging where available. For this analysis, we considered only cardioembolic, large vessel, and small vessel subtypes. For the cases that were classified as small vessel stroke under TOAST and had accompanying MRI, as well as all cases from UK Young Lacunar Stroke Study (DNAlacunar) which included only cases with the TOAST SVD subtype, we performed further characterization, as follows. All MRI scans were centrally reviewed by 1 physician (H.S.M.). The diagnosis of lacunar stroke was confirmed using the following criteria. Lacunar stroke was defined as a clinical lacunar syndrome,20 with an anatomically compatible lesion on MRI (subcortical infarct ≤15 mm in diameter). For MRIs performed in the acute phase, the diagnosis was made by an acute lacunar infarct on DWI. For scans not performed in the acute phase, the diagnosis was made by a lacunar syndrome in combination with a lacunar infarct visualized on T1 and fluid-attenuated inversion recovery as a cavitated lesion in an anatomically appropriate location. Exclusion criteria were as follows: stenosis >50% in the extra- or intracranial cerebral vessels; cardioembolic source of stroke, defined according to the TOAST criteria as high or moderate probability; subcortical infarct >15 mm in diameter, as these can be caused by embolic mechanisms (striatocapsular infarcts); any other specific cause of stroke (eg, lupus anticoagulant, cerebral vasculitis, dissection, monogenic forms of stroke, such as cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy). For each individual with a lacunar stroke, we characterized the degree of WMH using the semiquantitative Fazekas scale,²¹ which classifies individuals into 4 groups ranging from none (0) to severe (3). Based on this grading, we then divided the lacunar stroke cases into those with and without WMH: (1) no WMH—patients with only mild or absent leukoaraiosis (Fazekas grade 0 or 1) and (2) WMH-patients with moderate or severe leukoaraiosis (Fazekas grade ≥ 2).

Genetic Risk Score Analyses

For each of the 18 SNPs associated with WMH in community populations in a recent study (8 genome-wide and 10 with $P < 1 \times 10^{-5}$ in Europeans or overall),9 we generated an unweighted risk score for each individual in our data set by counting the number of risk alleles and summing across all SNPs. We used an unweighted approach, rather than an approach weighted on the log of the odds ratio (OR), as effect sizes were not reported for the published associations with WMH.9 Three of the SNPs (rs186314186, rs150695384, and rs117126031) were rare and not well imputed in our data set so were not included. Within each data set, we then converted each individual's risk score to a Z-score using the standard transformation. We then used logistic regression to estimate the influence of the risk score on each stroke outcome, including ancestry-informative principal components to control for population stratification and meta-analyzing the results using a fixed-effects inverse variance weighted approach. We tested for association of the genetic risk score with lacunar, cardioembolic, and large vessel stroke.

To investigate whether an association with lacunar stroke was independent of WMH, we performed the same analysis on 2 subgroups of lacunar stroke stratified on the presence of substantial WMH (Fazekas \geq 2). As the purpose of this analysis was to identify whether the observed association with lacunar stroke was independent of WMH, we did not perform the same analyses in the other subtypes. All ORs reported are per 1 SD change in the normally distributed risk score. We set the criteria for statistical significance at *P*<0.01, Bonferroni-correcting for the 5 tests.

Single SNP Analyses

In addition, we tested the association of each of the 15 available SNPs with lacunar stroke and the 2 subgroups based on the presence or absence of WMH. For each SNP we performed analyses separately in the 3 batches, including the first 10 ancestry-informative principal components in each analysis. We meta-analyzed the results using a fixed-effects inverse-variance weighted approach. We set the significance threshold at P<0.0011, correcting for the 15 SNPs in each of the 3 phenotypes (45 tests in total). All analyses were performed using the R statistical software.

Results

Cohort Characteristics

The final cohort consisted of 4176 stroke cases, including 1,373 lacunar stroke cases (mean age [SD]=60.0 [11.3] years; 68.0% men), subtyped into 568 with WMH (mean age [SD]=65.1 [10.9] years; 65.0% men) and 787 without WMH (mean age [SD]=56.0 [9.9] years; 70.2% men), 1,331 cardioembolic strokes (mean age [SD]=72.8 [10.6] years; 52.7% men), and 1,472 large vessel strokes (mean age [SD]=66.9 [11.1] years; 67.3% men) and 9,053 controls (mean age [SD]=58.4 [10.7] years [age not available in 2437 of WTCCC2-UK controls]; 52.2% men). Information on WMH volumes was not available in controls. As inclusion in the MRI-informed lacunar stroke analysis depended on the availability of an MRI and confirmation of a lacunar infarct, proportions of lacunar stroke cases varied greatly between studies (Figure 1).

Genetic Risk Score Analyses

A genetic risk score comprises 15 SNPs associated with WMH in community populations was significantly associated with lacunar stroke (OR [95% confidence interval [CI]=1.14 [1.06– 1.22]; P=0.0003; Figure 2). The association was slightly stronger, although not significantly so, in the group with WMH (OR [95% CI]=1.15 [1.05–1.26]; P=0.003) and slightly weaker and not reaching Bonferroni-corrected significance in the group without substantial WMH (OR [95% CI]=1.11 [1.02–1.21]; P=0.019). Conversely, the risk score was not associated with cardioembolic (OR [95% CI]=1.03 [0.97–1.09]; P=0.39) or large vessel stroke (OR [95% CI]=0.99 [0.93–1.04]; P=0.63).

Single SNP Analyses

No SNP reached the a priori significance threshold after Bonferroni correction (Table). Four SNPs (rs7909791[*SH3PXD2A*], OR [95% CI]=1.14 [1.03–1.27]; P=0.01; rs72934505[*NBEAL1*], OR [95% CI]=1.24 [1.07–1.45]; P=0.004; rs941898[*EVL*], OR [95% CI]=1.12 [1.01–1.25]; P=0.04; and rs9515201[*COL4A2*], OR [95% CI]=1.15 [1.03–1.27]; P=0.01) reached a nominal significance threshold in the all lacunar stroke analysis.

Discussion

We used a genetic risk score approach to determine whether genetic variants associated with WMH confer risk of lacunar stroke, and therefore whether WMH and lacunar stroke share pathogenesis. We found strong evidence that genetic variants associated with WMH in community populations also influence risk of lacunar stroke. This provides further evidence to support the long-held view that neuroimaging features

	DNA- Lacunar	GENESIS	WTCCC2 UK	WTCCC2 Germany	Milano	ASGC	LSS	
Total cases Total controls	930 970	351	2,374 5,175	I,174 797	372 407	1,162 1,249	459 455	
Toast CE Toast LVD Toast SVD	- - 930	80 64 82	460 498 474	330 346 106	65 74 25	240 421 310	157 70 55	
% of SVD with MRI <i>and</i> lacune confirmed	100	100	52.7	34.9	36.0	7.4	76.4	
% of MRI Confirmed SVD with WMH (Fazekas ≥2)	32.9	57.7	68.4	59.4	33.3	52.2	33.3	
Merge by similar genotyping array	r,	$\checkmark \downarrow$, /		_		
		DNA-Lacunar/ GENESIS]	MRI confirm stroke colla	ed lacunar aboration		LSS	TOTAL
Toast CE Toast LVD		80 64		1,094 1,338			157 70	1,331 1,472
MRI-confirmed SV - with WMH - without WMH	VD I	1,012 346 648		319 208 111	3		42 14 28	1,373 568 787
Controls		970		7,62	8		455	9,053

Figure 1. Cohort characteristics. ASGC indicates Australian Stroke Genetics Collaborative; CE, cardioembolic stroke; GENESIS, Genes and Ischaemic Stroke; LSS, Leuven Stroke Study; LVD, large vessel disease; MRI, magnetic resonance imaging; SVD, small vessel disease; WMH, white matter hyperintensities; and WTCCC2, Wellcome Trust Case Control Consortium 2.

of cerebral SVD share pathophysiology. When dividing our lacunar stroke population into those with moderate to severe WMH and those without, we found some evidence for association with both groups, although the association was marginally (and not significantly) stronger in the group with WMH and the association in the group without WMH did not reach Bonferroni-corrected significance. This suggests that variants influencing WMH confer risk of lacunar stroke even for lacunar strokes without substantial WMH. In addition, 2 of the SNPs, rs9515201[COL4A2] and rs2984613[PMF1-BGLAP], are also associated with intracerebral hemorrhage.^{10,22} This serves to emphasize that shared pathophysiological processes seem to underlie many of the clinical manifestations of cerebral SVD and suggests that a coordinated attempt to identify cerebral SVD associations will likely be fruitful. Four SNPs reached nominal significance for association with lacunar stroke (rs7909791[SH3PXD2A], rs72934505[NBEAL1], rs941898[EVL], and rs9515201[COL4A2]). With the exception of COL4A2, which has been linked to SVD, none of these loci have formerly been linked to ischemic stroke. As discussed above, similar arterial changes have been described in patients with lacunar stroke or WMH,5 including diffuse arteriosclerosis and a more focal microatheroma. Other studies have shown that mechanisms including blood-brain barrier dysfunction,²²⁻²⁴ and endothelial dysfunction,²⁵ are important in both.²³ As our results show a shared molecular basis to the 2 traits, they might suggest that these findings are because of the fact that WMH and lacunar stroke are outward manifestations of a shared underlying pathological process, namely, cerebral SVD.

In contrast, a genetic risk score comprising the same 15 SNPs was not associated with large vessel or cardioembolic stroke. Some studies have shown a relationship between subclinical atherosclerosis and WMH,^{23,24} whereas others have found an increased risk of all stroke in individuals with WMH.²⁵ Our results, in a well-characterized population, suggest that the relationship between WMH and ischemic stroke is limited to lacunar stroke. This finding might suggest that previously reported associations between nonlacunar strokes and WMH may be because of shared risk factors such as hypertension rather than shared pathogenesis.

This study has several strengths. The sample size was large and all lacunar strokes were confirmed by MRI, reducing the possibility of misclassification which might occur when using CT. In addition, the design of the study, which made use of genetic data, means that the results are less susceptible to the residual confounding and reverse causation that observational studies can suffer from, although other sources of confounding, such as technical artifacts, may arise. Similarly, this study has weaknesses. We were unable to evaluate 3 rare SNPs which were associated with WMH in the previous publication,⁹ which may have affected our results. Some lacunar infarcts were diagnosed as acute lesions on DWI, but others were diagnosed from MRI scans performed after the acute

Lacunar Stroke DNA-lacunar/GENESIS MCLSC LSS Overall, (p=0.0003)	₹ Ţ ↓		1.12 1.13 1.43 1.14	[1.03 , 1.23] [1.01 , 1.27] [1.02 , 2.01] [1.06 , 1.22]
Lacunar Stroke w/ WMH DNA-lacunar/GENESIS MCLSC LSS Overall, (p=0.003)			1.19 1.10 1.27 1.15	[1.05 , 1.35] [0.96 , 1.27] [0.73 , 2.20] [1.05 , 1.26]
Lacunar Stroke w/out WMH DNA-lacunar/GENESIS MCLSC LSS Overall, (p=0.019)	₽₽₽₽₽₽₽₽₽₽₽₽₽	• 1	1.08 1.17 1.53 1.11	[0.97 , 1.19] [0.96 , 1.42] [1.01 , 2.33] [1.02 , 1.21]
Large Vessel Stroke DNA-lacunar/GENESIS H MCLSC LSS Overall, (p=0.63)			0.81 0.99 1.09 0.99	[0.63 , 1.04] [0.93 , 1.05] [0.84 , 1.42] [0.93 , 1.04]
Cardioembolic Stroke DNA-lacunar/GENESIS MCLSC LSS Overall, (p=0.39)			0.92 1.05 0.95 1.03	[0.74 , 1.15] [0.98 , 1.12] [0.79 , 1.14] [0.97 , 1.09]
0.61	1.00	l 1.65 2	Т .72	
	Odds R	atio		

Figure 2. Association of white matter hyperintensities (WMH) Genetic Risk Score with ischemic stroke subtypes. GENESIS indicates Genes and Ischaemic Stroke; LSS, Leuven Stroke Study; and MCLSC, magnetic resonance imaging-confirmed lacunar stroke collaboration (Australian Stroke Genetics Collaborative [ASGC], Wellcome Trust Case Control Consortium 2 [WTCCC2-UK], Wellcome Trust Case Control Consortium 2 [WTCCC2]-Germany and Milano studies).

stroke phase as cavities on T1 or fluid-attenuated inversion recovery images. The inclusion of patients defined using these different radiological criteria may introduce a subtle bias, and it is possible that some of these cavities could have resulted from hemorrhage rather than ischemia. Although MRI was performed for all lacunar stroke subjects, it was not performed in all nonlacunar subtypes, which could lead to some degree of misclassification. In addition, controls were historical and MRI was not performed to rule out cerebrovascular disease.

Conclusions

Genetic factors that affect WMH are also associated with risk of lacunar stroke, but not other stroke subtypes. This sheds new light on processes that are implicated in lacunar stroke and provides further evidence that shared pathophysiological processes underlie different manifestations of SVD.

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Genetic Associations With White Matter Hyperintensities Confer Risk of Lacunar Stroke

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SUPPLEMENTAL MATERIAL

Genotyping and imputation

Cohorts genotyped on the Illumina 610K (Australian cases and controls, Italian cases), 660W (UK cases and controls, German cases and Italian controls) and 1M arrays (German controls) were treated as one batch. Quality control was performed on each dataset separately, as previously described. ¹⁻³ A consensus set of 381,428 SNPs was then identified that was consistent across the four populations and the four populations were merged for these SNPs. We then performed principal components analysis using EIGENSTRAT on an LD-pruned set of SNPs from the combined dataset removing any population outliers, ⁴ defined as greater than 6 standard deviations from the mean on the first 5 principal components. 37 individuals were removed in total (29 Australian, 8 Italian; 31 cases, 6 controls). The remaining individuals were then imputed to 1000 Genomes phase 3:⁵ SHAPEIT v2 was used to phase the haplotypes and IMPUTE v2.2.2 was used to perform the imputation.^{6,7}

The DNA-lacunar dataset was genotyped on the Illumina HumanExomeCore array. This array contains both exome content (~250,000 SNPs) and common tag SNPs (~250,000 SNPs) found on conventional GWAS arrays. Post-imputation, this array provides comparable coverage of common SNPs to larger arrays (MAF>5%, 78.2% $r^2 \ge 0.8$ compared to 86.6% for Illumina OmniExpress).⁸ SNPs were excluded with MAF<0.01, genotype missingness>3%, HWE p<1e-6 in controls, strand ambiguity (A/T or C/G) or evidence of differential missingness by case-control status (p<0.05). Individuals were excluded if they had missingness>3%, excess or reduced heterozygosity, showed evidence of relatedness with another individual (pi-hat>0.1875), or failed a "sex-check" in PLINK. EIGENSTRAT was used to remove non-caucasian individuals, and was then repeated to calculate ancestry-informative principal components.⁴ The remaining 269,691 autosomal SNPs and 2,603 individuals were then imputed to 1000 Genomes phase 3:⁵ SHAPEIT v2 was used to phase the haplotypes and IMPUTE v2.2.2 was used to perform the imputation.^{6,7}

The Leuven dataset (Leuven Stroke Study) was genotyped on the Illumina Omni 5M array. SNPs were excluded with MAF<0.01, genotype missingness>3%, HWE p<1e-6 in controls, strand ambiguity (A/T or C/G) or evidence of differential missingness by case-control status (p<0.05). Individuals were excluded if they had missingness>3%, excess or reduced heterozygosity, showed evidence of relatedness with another individual (pi-hat>0.1875), or failed a "sex-check" in PLINK. EIGENSTRAT was used to remove non-caucasian individuals. The remaining individuals were then imputed to 1000 Genomes phase 3:⁵ SHAPEIT v2 was used to phase the haplotypes and IMPUTE v2.2.2 was used to perform the imputation.^{6, 7}

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Appendix

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