## **CLINICAL AND POPULATION SCIENCES**

## Genome-Wide Association Study Meta-Analysis of Stroke in 22000 Individuals of African Descent Identifies Novel Associations With Stroke

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**BACKGROUND AND PURPOSE:** Stroke is a complex disease with multiple genetic and environmental risk factors. Blacks endure a nearly 2-fold greater risk of stroke and are 2× to 3× more likely to die from stroke than European Americans.

**METHODS**: The COMPASS (Consortium of Minority Population Genome-Wide Association Studies of Stroke) has conducted a genome-wide association meta-analysis of stroke in >22000 individuals of African ancestry (3734 cases, 18317 controls) from 13 cohorts.

**RESULTS:** In meta-analyses, we identified one single nucleotide polymorphism (rs55931441) near the *HNF1A* gene that reached genome-wide significance ( $P=4.62 \times 10^{-8}$ ) and an additional 29 variants with suggestive evidence of association ( $P<1\times10^{-6}$ ), representing 24 unique loci. For validation, a look-up analysis for a 100 kb region flanking the COMPASS single nucleotide polymorphism was performed in SiGN (Stroke Genetics Network) Europeans, SiGN Hispanics, and METASTROKE (Europeans). Using a stringent Bonferroni correction P value of  $2.08 \times 10^{-3}$  (0.05/24 unique loci), we were able to validate associations at the *HNF1A* locus in both SiGN ( $P=8.18\times10^{-4}$ ) and METASTROKE ( $P=1.72\times10^{-3}$ ) European populations. Overall, 16 of 24 loci showed evidence for validation across multiple populations. Previous studies have reported associations between variants in the *HNF1A* gene and lipids, C-reactive protein, and risk of coronary artery disease and stroke. Suggestive associations with variants in the *SFXN4* and *TMEM108* genes represent potential novel ischemic stroke loci.

**CONCLUSIONS:** These findings represent the most thorough investigation of genetic determinants of stroke in individuals of African descent, to date.

Key Words: brain ischemia Coronary artery disease genome-wide association study meta-analysis phenotype risk factors

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## Nonstandard Abbreviations and Acronyms

1000G	1000 genomes
ARIC	Atherosclerosis Risk in Communities
CHS	Cardiovascular Health Study
CIDR	Center for Inherited Disease Research
COMPASS	Consortium of Minority Population Genome-Wide Association Studies of Stroke
GEOS	Genetics of Early Onset Stroke
GWAS	genome-wide association study
HANDLS	Healthy Aging in Neighborhoods of Diversity across the Life Span
HNF1A	HNF1 homeobox A
ISGS	Ischemic Stroke Genetics Study
JHS	Jackson Heart Study
NINDS	National Institute of Neurological Disor- ders and Stroke
REGARDS	Reasons for Geographic and Racial Differences in Stroke
SiGN	Stroke Genetics Network
SIGNET	Sea Islands Genetics Network
SLESS	South London Ethnicity and Stroke Study
SNP	single nucleotide polymorphism
SWISS	Siblings with Ischemic Stroke Study
VISP	Vitamin Intervention for Stroke Prevention
WHI	Women's Health Initiative

Troke is the second leading cause of death worldwide and a leading cause of long-term disability in Uthe United States.<sup>1</sup> Stroke is a heterogeneous disease encompassing multiple subtypes with unique etiologies and risk factors.<sup>2</sup> Nearly 87% of the ≈795000 strokes that occur each year in the United States are ischemic.<sup>1</sup> Epidemiological studies suggest a substantial genetic component for stroke with overall heritability estimates of 38% for all ischemic strokes, and subtypespecific estimates of 20% to 25% for small vessel disease<sup>3</sup> and up to 40% for large-vessel disease.<sup>4</sup> Compared with European Americans, blacks have a nearly 2-fold greater risk of incident stroke, >2-fold increased risk of fatal stroke, strokes at younger ages, and higher frequency of poststroke disability.<sup>5,6</sup> Despite this disproportionate burden, few attempts to map stroke susceptibility loci have focused on individuals of African ancestry.<sup>7</sup> Recent genome-wide association studies (GWAS) have identified several stroke susceptibility loci8-14 primarily in individuals of European ancestry with little success replicating in non-European-ancestry populations<sup>7,13,15,16</sup> possibly due to differences in the genetic architecture of stroke among individuals of diverse ancestry.

This study represents a collective effort to investigate the genetic basis of stroke by mapping stroke susceptibility loci potentially unique to individuals of African ancestry. Using data obtained from the COMPASS (Consortium of Minority Population Genome-Wide Association Studies of Stroke), we expand upon our discovery GWAS meta-analysis of stroke in blacks<sup>7</sup> using 1000 genomes (1000G) imputed data in 22 000 individuals.

## **METHODS**

To minimize the possibility of unintentionally sharing information that can be used to reidentify private information, a subset of the data generated for this study are available at the database of Genotypes and Phenotypes (dbGaP) and can be accessed at https://www.ncbi.nlm.nih.gov/gap/.

## **Study Population**

COMPASS included a total of 22051 individuals of African descent with either a physician-adjudicated stroke (n=3734) or no history of stroke (n=18317; Table I in the Data Supplement) and genome-wide single nucleotide polymorphism (SNP) data. Participating studies include prospective cohorts (ARIC study [Atherosclerosis Risk in Communities],<sup>17</sup> CHS [Cardiovascular Health Study],18 JHS [Jackson Heart Study],19,20 the WHI [Women's Health Initiative],21]; casecontrol studies (INTERSTROKE,22 REGARDS [Reasons for Geographic and Racial Differences in Stroke],23 ISGS [Ischemic Stroke Genetics Study],24 VISP [Vitamin Intervention for Stroke Prevention],<sup>25,26</sup> SLESS [South London Ethnicity and Stroke Study],27 the GEOS Study [Genetics of Early Onset Stroke],28 the NINDS-SiGN [National Institute of Neurological Disorders and Stroke-Stroke Genetics Network],29 HANDLS [Healthy Aging in Neighborhoods of Diversity Across the Life Span]<sup>30</sup>); and an affected sib pair study-SWISS (Siblings With Ischemic Stroke Study).<sup>31</sup> Race/ethnicity-matched and sex-matched controls were randomly selected from HANDLS and used as controls in the analyses of SWISS, ISGS, and VISP, which lacked genotyped controls. All participants provided written, informed consent, and institutional review boards approved each of the respective studies/institutions.

## Outcomes

We defined stroke as a focal neurological deficit of presumed vascular cause with a sudden onset and lasting 24 hours or until death with clinical or radiological (computed tomography/magnetic resonance imaging) evidence with stroke diagnosis made when there is overwhelming clinical evidence in the absence of radiological evidence of a cerebral infarction. A lack of imaging data for all stroke cases does not increase the likelihood of false positives in our study. The cohort studies only considered first (incident) clinically validated ischemic strokes. Individuals with a baseline history of ischemic or hemorrhagic stroke were excluded.

## Genotype Data

All studies imputed SNPs using 1000G Phase I Version 3 Haplotypes, except SLESS and WHI, which used 1000G Phase III data (1KGp3) reference populations. We excluded

<b>Table</b>	able 1. COMPASS Ischemic Stroke Suggestive and Genome-Wide Significant Inverse Variance Weighted Associations										ns	
Chr	Position*	Gene	SNP	Alleles (Coded/ Noncoded)	Beta	SE	Odds Ratio (Cl)	Inverse Variance Weighted <i>P</i> Value	Direction	Het <i>P</i> Value	Sample Size	No. of Studies
1	112853017	CTTNBP2NL (nearest)	rs114947355	T/C	0.44	0.0902	1.56 (1.42–1.70)	9.05×10 <sup>-07</sup>	????-+?+???	0.1382	12610	3
1	112857084	CTTNBP2NL (nearest)	rs147779128	A/T	-0.46	0.0945	0.63	9.61×10 <sup>-07</sup>	?????~?~????	0.9293	9637	2
2	4083658	NPM1P48 (nearest)	rs142655108	A/C	0.58	0.1089	1.79 (1.60–1.99)	9.52×10 <sup>-08</sup>	?????+?+????	0.2834	9637	2
2	198551159	RFTN2 and MARS2 (nearest)	rs115670077	T/G	0.35	0.072	1.43 (1.33–1.53)	8.48×10 <sup>-07</sup>	+?++++?+???	0.5735	16540	6
3	124048486	KALRN	rs72976591	A/C	0.17	0.0342	1.18 (1.14–1.22)	9.19×10 <sup>-07</sup>	+++++++++++++++++++++++++++++++++++++++	0.5356	22018	11
3	133101791	TMEM108	rs113509723	-/AA	0.45	0.0841	1.58 (1.45–1.71)	6.46×10 <sup>-08</sup>	?????+?+???	0.2014	9637	2
3	153125290	AK092619 (nearest)	rs184221467	A/G	0.62	0.1246	1.85 (1.63–2.10)	7.86×10 <sup>-07</sup>	?????+?+???	0.468	9637	2
4	99435032	TSPAN5	rs138134155	A/G	0.36	0.0705	1.43 (1.33–1.53)	3.94×10 <sup>-07</sup>	+?+++?++??	0.9442	18531	7
5	101123995	OR7H2P (nearest)	rs77460585	A/G	0.59	0.1165	1.80 (1.60–2.02)	4.36×10 <sup>-07</sup>	????-??+???	0.004981	10940	2
5	150981704	FAT2 and SPARC (nearest)	rs114527838	A/G	-0.28	0.055	0.76 (0.72–0.80)	5.55×10 <sup>-07</sup>	-???	0.7033	19032	8
6	97345991	<i>KLHL32</i> and <i>NDUFA4</i> (nearest)	rs146522546	-/CT	-0.45	0.0876	0.64 (0.58–0.69)	2.22×10 <sup>-07</sup>	????-????	0.3829	13353	4
7	83432409	SEMA3A	rs6967981	T/G	0.15	0.0296	1.16 (1.12–1.19)	7.57×10 <sup>-07</sup>	+-++++++++-	0.1685	21970	11
8	1572874	DLGAP2	rs112455974	A/C	0.68	0.1336	1.97 (1.72–2.25)	3.77×10 <sup>-07</sup>	????+??+???	0.7366	10949	2
9	72475192	C9orf135	rs565295967	T/C	0.62	0.1199	1.86 (1.65–2.09)	2.41×10 <sup>-07</sup>	?????+?+???	0.1048	9637	2
10	53545098	PRKG1	rs140164788	T/C	0.52	0.1019	1.68 (1.52–1.86)	3.37×10 <sup>-07</sup>	????++?+???	0.7146	12618	3
10	53547264	PRKG1	rs74469072	T/G	0.52	0.1018	1.68 (1.52–1.86)	3.50×10 <sup>-07</sup>	????++?+???	0.7169	12618	3
10	120907173	SFXN4	rs150807690	—/G	-0.20	0.0378	0.82 (0.79–0.85)	9.67×10 <sup>-08</sup>	?-??	0.3014	18180	8
11	11360296	GALNT18	rs115825287	T/C	0.35	0.0696	1.43 (1.33–1.53)	3.60×10 <sup>-07</sup>	??++++?+???	0.6076	15673	5
11	75683895	UVRAG	rs368167310	T/C	-0.55	0.1085	0.58 (0.52–0.65)	4.87×10 <sup>-07</sup>	?????-?-????	0.8172	9637	2
12	29288407	FAR2 (nearest)	rs113025543	A/T	-0.27	0.0551	0.76 (0.72–0.81)	9.23×10 <sup>-07</sup>	-+?	0.7896	20224	10
12	29292793	FAR2 (nearest)	rs142100833	C/G	0.24	0.0488	1.27 (1.21–1.34)	8.65×10 <sup>-07</sup>	+-++++++-?	0.4482	20119	10
12	29341407	FAR2	_	-/??	0.65	0.1272	1.91 (1.68–2.17)	3.79×10 <sup>-07</sup>	???++???+??	0.9784	5542	3
12	119502791	SRRM4	rs531465435	—/C	0.59	0.1162	1.81 (1.61–2.03)	3.39×10 <sup>-07</sup>	?????+?+????	0.5809	9637	2
12	119542751	SRRM4	rs192977447	A/T	0.43	0.0816	1.53 (1.41–1.66)	1.80×10 <sup>-07</sup>	???+++?++??	0.1962	15333	5
12†	121415209	HNF1A (nearest)	rs55931441	A/G	0.52	0.0947	1.68 (1.53–1.84)	4.62×10 <sup>-08</sup>	?????+?+???	0.4599	9637	2
14	93788855	BTBD7	rs113949028	—/G	0.20	0.0396	1.22 (1.17–1.27)	5.44×10 <sup>-07</sup>	?+?+++?++++	0.948	18255	8
18	68475060	GTSCR1 (nearest)	rs181095590	A/G	0.58	0.1138	1.78 (1.59–2.00)	3.90×10 <sup>-07</sup>	?????+?+????	0.4538	9637	2
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#### (Continued)

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#### Table 1. Continued

Chr	Position*	Gene	SNP	Alleles (Coded/ Noncoded)	Beta	SE	Odds Ratio (CI)	Inverse Variance Weighted <i>P</i> Value	Direction	Het <i>P</i> Value	Sample Size	No. of Studies
19	29710081	UQCRFS1 (nearest)	rs73923591	A/G	0.27	0.0548	1.31 (1.24–1.39)	6.18×10 <sup>-07</sup>	++++++++?	0.8774	20246	10
21	36442465	RUNX1	rs116262092	A/T	-0.58	0.1174	0.56 (0.50–0.63)	7.04×10 <sup>-07</sup>	\$5555-555	0.9789	12581	3
21	36443919	RUNX1	rs147867382	C/G	-0.58	0.1174	0.56 (0.50–0.63)	7.95×10 <sup>-07</sup>	?????-????	0.9792	12579	3

Direction indicates the direction of the effect size: negative (–), neutral/unknown (/?), and positive (+) for each contributing cohort/population. Chr indicates chromosomes; COMPASS, Consortium of Minority Population Genome-Wide Association Studies of Stroke; Het, heterogeneity; and SNP, single nucleotide polymorphism. \*Chr position based on human genome (GRCh37/hg19).

†Genome-wide significance ( $P < 5 \times 10^{-8}$ ).



Figure. LocusZoom plots, with linkage disequilibrium based on hg19/1000 Genomes Nov 2014 AFR, depicting the top ( $P=10^{-8}$ ) 3 associations with ischemic stroke in COMPASS (Consortium of Minority Population Genome-Wide Association Studies of Stroke) individuals of African descent.

**A**, *HNF1A* (rs55931441) chromosome (chr) 12 locus; (**B**) *TMEM108* (rs113509723) chr3 locus; (**C**) chr2 (rs142655108) locus nearest *NPM1P48*. SNP indicates single nucleotide polymorphism.

SNPs if they had invalid or missing alleles, *P* values, or  $\beta$  values; had minor allele frequencies <1%; imputation quality (*r*<sup>2</sup>) <0.3; or were located on sex chromosomes. We analyzed SNPs available in ≥2 studies, for a total of ≈16.9 million SNPs. The Data Supplement contains study-specific details about design, stroke definition, adjudication procedures, and genotyping.

### Analysis

We used logistic regression (additive genetic model) analyses with a count of variant alleles (0, 1, or 2) for each genotyped SNP or allelic dose for imputed SNPs. To control for potential population stratification, we included estimated study-specific principal components of global ancestry as covariates. As

Chr	Unique Locus	Top SiGN European SNP	Alleles	Z Score	P Value	Direction	Top SiGN Hispanic SNP
1	CTTNBP2NL (nearest)	rs186896391	C/A	-3.28	0.0010*	+++	rs3121986
2	NPM1P48 (nearest)	2-4077298 (rs527602504)	TC/T	2.56	0.0104	+	rs60037207
2	RFTN2 and MARS2 (nearest)	2-198592085 (rs543821034)	C/T	2.98	0.0029	+	rs150235598
3	KALRN	rs2034173	T/C	2.99	0.0027	++	rs185731506
3	TMEM108	rs13087036	C/A	-2.52	0.0116	++	rs139695007
3	AK092619 (nearest)	rs183598421	T/C	-2.36	0.0185		rs200248409
4	TSPAN5	rs28392914	T/G	-3.16	0.0016*	++	rs1045655
5	OR7H2P (nearest)	rs139061870	GT/G	2.80	0.0052	+	rs73776672
5	FAT2 and SPARC (nearest)	rs141575897	G/A	-3.03	0.0024	+	rs80009114
6	KLHL32 and NDUFA4 (nearest)	rs200056339	C/CA	-2.68	0.0074		rs78235656
7	SEMA3A	rs151172774	T/C	2.76	0.0058	+++++++++++++++++++++++++++++++++++++++	rs6955094
8	DLGAP2	rs117175403	G/A	2.79	0.0053	-++++++.	rs184526444
9	C9orf135	rs56179412	C/T	-2.13	0.0330	+	rs77797545
10	PRKG1	rs10999787	C/A	-2.70	0.0069	++	rs10998992
10	SFXN4	rs143931152	T/G	-3.64	0.0003*		rs56095167
11	GALNT18	rs117835740	C/T	-2.45	0.0142		rs11021735
11	UVRAG	11-75761242 (rs565239444)	T/G	-2.76	0.0058		rs138825035
12	FAR2 (nearest)	rs151183596	T/A	-2.70	0.0070	-++	rs141911197
12	SRRM4	rs61937966	C/T	3.37	0.0007*	+++++++++++++++++++++++++++++++++++++++	rs4767761
12	HNF1A (nearest)*	rs182546302	T/A	-3.35	0.0008*	-+++.	rs80019595
14	BTBD7	rs112848587	C/T	-2.19	0.0284		rs76789831
18	GTSCR1 (nearest)	rs11151610	T/C	-3.27	0.0011*		rs75968601
19	UQCRFS1 (nearest)	rs148613358	T/C	3.22	0.0013*	+++.	rs12608817
21	RUNX1	rs7280028	T/C	-3.42	0.0006*		rs9981811

Table 2.	Genome-Wide and Suggestive COMPASS Associations With Look-Ups in European and Hispanic Populations From
SiGN and	I METASTROKE

(Continued)

appropriate, we adjusted models for age, sex, and study site. We combined study-specific results in a fixed-effects meta-analyses with inverse variance weighting using METAL.<sup>32</sup> We also performed sample size weighted meta-analysis as an alternative approach to inverse variance weighting (Table II in the Data Supplement). We set a genome-wide significance (discovery) threshold of  $P < 5 \times 10^{-8}$  but investigated all SNPs with  $P < 10^{-6}$ .

## Validation of COMPASS Findings

Due to the absence of a comparable and adequately powered cohort of blacks with GWAS and adjudicated stroke data, we performed a look-up of COMPASS SNPs with  $P < 10^{-6}$  in the SiGN European and Hispanic ischemic stroke populations and METASTROKE total ischemic stroke populations (Table III in the Data Supplement). Additional METASTROKE subtype (cardioembolic, large-vessel, and small vessel) specific look-up analyses were performed to further validate these findings. Given the known differences in linkage disequilibrium patterns between populations of European and African ancestry, we

expanded the region of interest for each locus to include available SNPs±100 kb of the index COMPASS SNPs as previously described<sup>7</sup> applying a Bonferroni correction to account for the number of loci tested.

## RESULTS

### **Discovery of Stroke-Associated Loci**

Using inverse variance weighting meta-analyses (Table 1), we identified one genome-wide significant association ( $P < 5 \times 10^{-8}$ ) and an additional 29 variants with suggestive evidence of association ( $P < 1 \times 10^{-6}$ ), representing 24 unique loci in total. The genome-wide significant association was detected upstream of the HNF1 homeobox A (*HNF1A*) gene on chromosome 12 (rs55931441;  $P = 4.62 \times 10^{-8}$ , odds ratio, 1.68; Figure [A]).

#### Table 2. Continued

Alleles	Z Score	P Value	Direction	METASTROKE Top SNP	Alleles	Effect	<i>P</i> Value	Direction	
A/G	-2.79	0.0052	_	rs10158830	C/G	0.073	0.0019*	++++++	
T/C	-2.21	0.0268	_	rs114152357	A/T	-0.186	0.0048	+-++	
G/A	-2.74	0.0061	-	rs191948652	A/T	0.513	0.005	+-+???+++?+?	
C/G	-3.11	0.0019*	_	rs73188175	T/C	0.300	0.0019*	+++++++++++++++++++++++++++++++++++++	
G/C	3.09	0.0020*	+	rs2699882	A/G	0.053	0.0096	+-+++++++	
GT/G	-2.86	0.0043	-	rs7427054	T/C	0.093	0.0015*	++++++++-	
G/C	-2.87	0.0041	-	rs12509107	A/G	-0.445	0.0168		
T/C	-3.43	0.0006*	-	rs62386289	T/C	-0.117	0.0039	-++++	
A/G	2.53	0.0113	+	rs6579892	A/T	0.075	0.00095*	++++++++++++	
G/A	-2.77	0.0057	-	rs117804808	T/C	0.250	0.0099	+++-+++++++++++++++++++++++++++++++++++	
A/G	3.18	0.0015*	+	rs150770834	A/G	0.494	0.0108	-????++++++	
A/T	-2.90	0.0037	-	rs11998452	A/G	-0.218	0.0021	+-+?-?	
A/G	2.29	0.0220	+	rs143862820	T/C	0.289	0.0055	?++-?+++-+??	
C/T	-2.81	0.0049	-	rs192204676	A/G	0.332	0.016	+??++++++	
G/A	-3.21	0.0013*	_	rs188855777	T/C	-0.653	0.0032	-??-?-?-?????	
C/T	2.90	0.0037	+	rs4909989	A/G	-0.080	0.0033	++-	
A/G	-3.39	0.0007*	_	rs139079454	T/C	0.233	0.0043	+++?++++-+++	
T/G	-3.50	0.0005*	-	rs12311115	A/G	-0.119	0.00031*	++	
A/G	-3.40	0.0007	-	rs78381318	A/G	0.194	0.000013*	++-++++++++++++++++++++++++++++++++++++	
C/T	-2.62	0.0087	_	rs117548270	A/G	-0.312	0.0017*	+?	
C/G	-2.77	0.0057	-	rs111650311	T/C	0.072	0.0228	++++-+++++?+	
C/T	2.98	0.0029	+	rs146227033	C/G	-0.245	0.00068*	?+	
C/A	-3.12	0.0018*	-	rs2160742	A/G	0.074	0.0047	++-++++-+++	
G/A	2.92	0.0035	+	rs2247822	T/C	0.071	0.00055*	++++++++-	

Chr indicates chromosomes; COMPASS, Consortium of Minority Population Genome-Wide Association Studies of Stroke; SiGN, Stroke Genetics Network; and SNP, single nucleotide polymorphism.

\*Significance for replication P<2.08x10<sup>-3</sup>).

# Validation of COMPASS SNPs in SiGN and METASTROKE

Expanding to the flanking regions and using a stringent Bonferroni correction of  $\alpha$ =2.08×10<sup>-3</sup> for replication (0.05/24 unique loci), our most significant locus, *HNF1A*, was validated in both SiGN and METASTROKE European-ancestry cohorts and approached significance in SiGN Hispanics (Figure I in the Data Supplement). Overall, 16 of 24 loci showed evidence for validation across multiple populations (Table 2).

Likely due to the inclusion of ischemic stroke cases only, we were not able to replicate the novel association for rs4471613, which was associated with total (ischemic and hemorrhagic) stroke in our prior COMPASS HapMap imputation report (inverse variance weighting P=0.85).<sup>7</sup> Additionally, we found no evidence of replication for loci previously associated with stroke in European-Ancestry populations (*P* ranging from 0.02 to 0.95; Tables IV and V in the Data Supplement).

## DISCUSSION

This new COMPASS meta-analysis of ischemic stroke only identified 24 unique loci with suggestive (n=23) or genome-wide (n=1) evidence for association with ischemic stroke. The most significantly associated *HNF1A* variant, rs55931441 (G/A), is monomorphic in European populations (G allele present only), with a 2% minor allele frequency (allele A) reported in sub-Saharan and 1000G African populations, and 3.8% frequency in **CLINICAL AND POPULATION** 

COMPASS. This SNP was present in the only 2 studies imputed to 1000G Phase III (WHI and SLESS). Collectively, WHI and SLESS account for 9637 subjects (1147 stroke cases and 8490 controls). We were unable to assess the association for rs55931441 directly in our cross-ethnic look-up; however, SNPs in a 100 kb flanking region were significant (Figure I in the Data Supplement) in SiGN Europeans (top SNP rs182546302; P=8.18×10<sup>-4</sup>), METASTROKE ischemic stroke phenotype (top SNP rs117548270; P=1.72×10<sup>-3</sup>), and METASTROKE cardioembolic stroke phenotype (top SNP rs184865012; P=9.98×10<sup>-4</sup>), whereas SNP rs80019595 approached significance ( $P=8.74\times10^{-3}$ ) in the SiGN Hispanic cohort. Previous studies have reported associations between variants in HNF1A and lipids,<sup>33</sup> C-reactive protein,<sup>34,35</sup> and risk of coronary artery disease and stroke.33,35 Taken together, these findings may provide greater insight regarding subtypespecific influences and potential mechanism of HNF1A variants in stroke risk.

Three additional variants reached suggestive associations at the  $P \le 10^{-8}$  level (rs113509723 in TMEM108 (Figure [B]); rs142655108 near *NPM1P48* (Figure [C]); rs150807690 in SFXN4). The NPM1P48 locus showed no evidence for replication in the cross-ethnic look-up, whereas TMEM108 was replicated in SiGN Hispanics only (top SiGN Hispanic SNP rs139695007; P=0.002). The SFXN4 SNP, rs150807690, is a G insertion (-/G)with a 22% minor allele frequency (G insertion) in the 1000G African population and 24% frequency in COM-PASS. Variant rs150807690 did not replicate in SiGN Hispanic (P=0.796) or SiGN Europeans (P=0.696) analyses and was not present in the METASTROKE look-up; however, nearby SNPs with evidence of replication in a 100 kb flanking region were detected in SiGN Europeans (top SNP rs143931152; P=2.68×10<sup>-4</sup>) and SiGN Hispanics (top SNP rs56095167; P=1.31×10<sup>-3</sup>), located 35540 bp and 97388 bp from the indexed COMPASS variant, respectively. The SFXN4 gene has not been previously implicated in stroke. The protein encoded by SFXN4 is critical for mitochondrial respiration and erythropoiesis.36,37 Recent clinical trials suggest that erythropoiesis-stimulating agents effectively treat anemia associated with chronic kidney disease but increase the risk of stroke possibly due to hyperviscosity.<sup>38</sup>

Of the 23 loci with suggestive association in COM-PASS, 15 showed evidence for replication in  $\geq$ 1 look-up analysis. One locus was replicated in SiGN Europeans only, four loci were replicated in SiGN Hispanics only, 2 loci were replicated in METASTROKE ischemic stroke only, whereas 8 loci had evidence for replication in  $\geq$ 2 look-ups. Two loci, *SFXN4 and UQCRFS1*, were replicated in both the SiGN Europeans and Hispanics, 2 loci were replicated in SiGN Hispanics and METASTROKE ischemic stroke (*KALRN* and *FAR2*), and 3 loci were replicated in SiGN Europeans and METASTROKE ischemic stroke (*CTTNBP2L*, *GTSCR1*, and *RUNX1*). Most notably, one locus (*SRRM4*) was replicated in all 3 look-ups. Evidence for association across multiple ethnicities might indicate stroke susceptibility loci with a global impact. For example, the *KALRN* locus which was replicated in SiGN Hispanics and METASTROKE has been implicated in coronary artery disease risk across multiple populations<sup>39–41</sup> and was recently associated with ischemic stroke and lacunar stroke in a Han Chinese population.<sup>42</sup> Although the *SRRM4* locus, which was replicated in all 3 look-ups, has not previously been implicated in stroke, the gene is important for neurogenesis<sup>43</sup> and has shown associations with neurological conditions including Alzheimer disease<sup>44</sup> and epilepsy.<sup>45</sup>

Although this effort represents the largest stroke GWAS meta-analysis in individuals of African descent, the modest sample size of 3734 stroke cases limits our power to detect associations for variants with minor allele frequencies of ≤3%. Only 2 cohorts used the most recent imputation panel limiting our ability, and thus power, to detect novel variants only present in 1000G Phase III and not 1000G Phase I Version 3. Furthermore, individuals of African descent experience ischemic strokes of small vessel origin more frequently. Therefore, due to the increased genetic diversity of this COMPASS population combined with the greater prevalence of small vessel stroke, we are not surprised at a lack of validation of previous European-ancestry associations. Failure to replicate associations across ethnicities is a common occurrence in genetic studies of various diseases and, therefore, does not threaten the validity of our current study. Moreover, the lack of availability of an adequate replication cohort consisting of individuals of African descent suffering a stroke that have genomewide SNP genotype data remains a substantial global challenge. Likewise, due to smaller linkage disequilibrium blocks and increased genetic diversity in populations of African descent, larger sample sizes would help alleviate limitations of statistical power, challenges associated with imputing genotypes, and allow for more detailed stroke subtype analyses. A recent analysis showed that although the number of GWAS conducted as of 2016 has increased >6-fold since 2009, African descent participants increased by only 2.5%.46 Therefore, our study will help advance precision medicine applications by identifying genetic loci (and subsequent polygenic risk scores) for stroke prediction and risk stratification in diverse populations.

### SUMMARY

Despite its limitations, genetic studies, such as COM-PASS, that include minority populations have the huge potential to provide insight into the mechanisms underlying stroke disparities, such as the more than doubled incidence and mortality rates and younger age of onset for stroke observed in blacks.<sup>5,47</sup> Our study identified novel associations for stroke that might not otherwise be detected in primarily European cohort studies. Collectively, this highlights the critical nature and importance of genetic studies in a more diverse population with a high stroke burden, such as was the case in this study.

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