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Genome-wide association study of early ischemic stroke risk in Brazilian Individuals with Sickle Cell Disease implicates *ADAMTS2*, *CDK18* and uncovers novel loci

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Abstract

Ischemic stroke is a common complication of sickle cell disease (SCD) and without intervention can affect 11% of children with SCD before the age of 20. Within the Trans-Omics for

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Precision Medicine (TOPMed), a genome-wide association study (GWAS) of ischemic stroke was performed on 1,333 individuals with SCD from Brazil (178 cases, 1,155 controls). Via a novel Cox proportional hazards analysis, we searched for variants associated with ischemic stroke occurring at younger ages. Variants at genome wide significance ($P < 5 \times 10^{-8}$) include two near genes previously linked to non-SCD early onset stroke (<65 years): *ADAMTS2* (rs147625068, $P=3.70 \times 10^{-9}$) and *CDK18* (rs12144136, $P=2.38 \times 10^{-9}$). Meta-analysis which included the independent SCD cohorts Walk-PHaSST and PUSH exhibited consistent association for variants rs1209987 near gene *TBC1D32* (P = 3.36×10^{-10}), rs188599171 near *CUX1* ($P=5.89 \times 10^{-11}$), rs77900855 near *BTG1* (P= 4.66×10^{-8}), and rs141674494 near *VPS13C* (1.68×10^{-9}). Finding from this study support a multi-variant model of early ischemic stroke risk and possibly a shared genetic architecture between SCD individuals and non-SCD individuals <65 years.

Keywords

GWAS; sickle cell disease; Brazil; stroke; ischemic stroke

INTRODUCTION

Ischemic stroke is one of the most common comorbidities of sickle cell disease (SCD), occurring 20 times more frequently than seen in non-SCD children^{1,2}. Without early preventative therapy, 11% of children with SCD experience at least one overt ischemic stroke before the age of 20³, with recurrence ranging from 60–92%⁴ and mortality of roughly 5%⁵. Ischemic stroke is more common than hemorrhagic stroke in children, with hemorrhagic stroke more common in ages 20–35 years. The pathophysiology of ischemic stroke within SCD is complex¹, and the current standard for risk assessment, transcranial doppler ultrasonography (TCD) of cerebral vessels, is difficult to perform on children under age 3. Also, some individuals with normal TCD still experience stoke^{3,6,7}.

Genetic risk of stroke has been investigated in SCD and non-SCD populations. Within SCD, stroke risk is highest in *HbSS* homozygotes compared to other SCD genotypes, whereas the coinheritance of alpha-thalassemia is protective against stroke¹. There is mounting evidence for the existence of other heritable genetic risk factors for stroke.^{8–15} A GWAS of 677 African Americans with SCD discovered two nonsynonymous variants in the genes *GOLGB1* and *ENPP1* associated with protection from stroke¹³. The same variant in *ENPP1* was confirmed within a Brazilian SCD cohort, however *GOLGB1* was not replicated¹⁵. Recently, *APOE* variants were also shown to be associated with ischemic stroke in children with SCD ages four and younger¹⁶.

Investigating genetic risk within non-SCD stroke may provide additional insight into the similarity or differences compared to SCD. Within non-SCD pediatric ischemic stroke studies, risk variants near genes *ADAMTS2, ADAMTS12*, and *ADAMTS13* haven been discovered, and both *ADAMTS2* and *ADAMTS12* have been replicated^{17,18}. In separate studies, variants near *CDK18*¹⁹, *HABP2*²⁰, were linked to early onset ischemic stroke but have not been replicated. Gaps in our knowledge of the genetic contributors to ischemic stroke risk within SCD remain. To address this, we conducted the largest genome-wide

association study of stroke in individuals with sickle cell disease (N=1,333). This was conducted within the Brazil Sickle Cell Disease Cohort Study as part of the Recipient Epidemiology Donor Evaluation Study III (REDS-III)²¹.

METHODS

Two novel approaches were implemented in this study. Variants were measured via whole genome sequencing, avoiding the need to impute on reference panels which may have little to no overlap in ancestry with the study cohort. In addition, a Cox proportional hazards model was used to estimate differences in time to stroke, allowing for the discovery of genetic variants associated with earlier stroke events.

Study Population

The Brazil SCD Cohort Study is part of the National Institutes of Health, National Heart Lung and Blood Institute (NHLBI) REDS-III program.²² The Brazil National Research Ethics Commission, local ethical committees at each participating center and the Institutional Review Boards at University of California, San Francisco (UCSF) and the REDS-III data coordinating center, Research Triangle Institute, International (RTI) all reviewed and approved the study.

2,793 sickle cell disease individuals were enrolled at 6 different centers in Brazil: Hemominas Belo Horizonte, Juiz de Fora, and Montes Claros, Hemorio, Rio de Janeiro, Instituto de Tratamento do Câncer Infantil, Sao Paulo, and Hemope, Recife. Enrollment included interviews, medical abstraction, and blood collection. Written informed consent was obtained from participants 18 years or from guardians of younger patients, and assent was obtained for children aged 7 to 17. Medical records were abstracted for clinical history using standardized definitions²³. Participants were routine patients at these centers, and relatively complete medical histories were available, including acute complications. This included a history of ischemic stroke defined as an acute neurological syndrome resulting from impaired cerebral blood flow without evidence of hemorrhage. Diagnostic criteria included either a magnetic resonance image (MRI) or a computerized tomography (CT) scan showing an infarctive central nervous system event consistent with symptoms and signs or diagnosis based on examination and clinical history with neurologic symptoms/signs lasting >24 hours. Controls were defined as individuals with SCD who had no history of stroke. While MRI/CT was available to confirm cases, current guidelines for care of SCD does not include MRI/CT when neurological symptoms are absent.

DNA collection and sequencing

Whole blood was collected in ethylenediaminetetraacetic acid (EDTA), and DNA was extracted from the buffy coat via ethanol precipitation. Purified DNA was quantified with RT-PCR, and normalized to 10 ng/uL. SCD genotypes were confirmed using allele-specific pyrosequencing (Qiagen, Hilden, Germany)²⁴ and Sanger sequencing of exons 1 and 2 of gene *HBB* if the pyrosequencing results conflicted with medical records. Additional sequencing of *HBB* exon 3, introns, and promoter was performed for samples with

unresolved genotypes at the Hemoglobinopathy Reference Laboratory at UCSF Benioff Children's Hospital Oakland.

Whole genome sequencing was conducted within the NHLBI Trans-Omics for Precision Medicine (TOPMed) program (freeze 6a)²⁵. Average sequencing depth was 38 fold, and variants were called jointly across roughly 140,000 samples, including the N=1,333 individuals in the current study, using the GotCloud pipeline.²⁵ The list of roughly 18 million variants were filtered to include only those at observed minor allele frequency >1%, resulting in a final list of 14.3 million variant autosomal sites.

Exclusion Criteria

Individuals were excluded from this analysis for the following reasons: not having TOPMed DNA sequence data, being a duplicate sample, inability to link DNA to clinical records, missing age or stroke status, and not being an *HbSS* homozygote. Samples were also removed if they were clear outliers along principal components 1–3 of the genotype data. Finally, we also excluded individuals with no history of stroke who were either undergoing chronic transfusion therapy (CTT) or had abnormal transcranial doppler (TCD) readings (Supplemental Fig 1). This was done to ensure a more homogenous population included in the study.

Statistical Analyses

A Cox proportional hazards model was used to measure single nucleotide polymorphisms (SNP) associations. Age at stroke was used for cases, and age at enrollment was right censored for controls without a history of ischemic stroke. Principal components (PC) were calculated from LD pruned genotypes²⁶ using the entire freeze6a TOPMed cohort and was then was subset to the N=1,333 participants in the study. Statistical tests were performed using the R package *survival*²⁷ with the following model:

 $\lambda(t|Stroke_i) = \lambda_0(t)\exp(\beta_1 SNP_i + \beta_2 PC1_i \dots + PC10_i)$

Where *t* is time in years to first stroke (or years to enrollment for controls), *PC* is the principal component eigenvector of the observed genotype covariance matrix, and *SNP* represents the genotype of the individual. Genotypes were converted to integers prior to populating the model in the following way: 0 = homozygous reference; 1 =heterozygous reference/variant; 2 = homozygous variant. Genome-wide significance was defined as $P < 5 \times 10^{-8}$. Conditional regression was performed on candidate 5Mb genomic regions surrounding the lead SNPs defined by lowest *P*-value. No independent loci within the candidate regions were observed. Code for this analysis is available online: https:// github.com/earleyej/cox-gwas.

All genotype-level analyses were performed in a HIPPA compliant environment on DNAnexus (https://www.dnanexus.com/). Annotated views of genomic candidate regions were created with LocusZoom²⁸ using the full (ALL) 1000 genomes reference for linkage disequilibrium (LD) estimates. Tests on GWAS summary statistics, including gene-level association and enrichment tests were performed using the online portal Functional Mapping

and Annotation of GWAS (FUMA²⁹). Gene-level association was performed on the full 14 million *P*-values using Multi-marker Analysis of GenoMic Annotation (MAGMA)³⁰ using default settings. Gene set pathway analysis was performed on the Gene Ontology (GO) database (N=9,988). Colocalization analysis was performed with the R package *coloc*³¹ on 1Mb regions surrounding each independent genome-wide significant SNP with the lowest *P*-value for that region (aka 'lead' SNP). Additional gene set enrichment was performed on the GWAS Catalog³² of genes using hypergeometric tests and corrected for multiple hypothesis testing using Benjamini-Hochberg FDR³³. Candidate genes were defined as being within 500Kb of a lead SNP, or if no genes were present within 500Kb, then the most proximal gene was used for that region. Meta-analysis of candidate SNPs with Walk-PHaSST and PUSH cohorts was performed using Fisher's method.

Multi-variant survival analysis

Individuals were categorized reflecting the number of lead SNPs they carried. All categories were required to have at least 10 participants, for a total of 5 categories (0, 1, 2, 3, and 4). These were used as categorical variables in a univariate survival analysis. *P*-values were calculated using Gehan-Wilcoxon³⁴, a non-parametric test which has more sensitivity in detecting differences between groups where the hazard ratio is higher at earlier timepoints.

RESULTS

Cohort characteristics

Of the 2,793 SCD individuals enrolled in the REDS-III Brazil SCD Cohort, 1,460 were excluded, with a final set of 1,333 individuals (Supplemental Figure 1). Younger age was strongly associated with ischemic stroke (OR=0.9; CI=0.89–0.92; P<0.001; Figure 1A) but not sex (male risk OR=0.9; CI, 0.65–1.24, P=0.5) or Hemocenter (OR=0.98; CI, 0.89–1.08; P=0.7). Many patients reported taking hydroxyurea (HU); however, the majority of participants initiated treatment after their first stroke and thus we did not consider HU in our analysis for time to first ischemic stroke. Genome-wide association was conducted on *HbSS* homozygotes only (N=1,333). Within this set were 178 (13.4%) individuals who experienced ischemic stroke (Table 1) with a median age at stroke of 9.5 years (Table 1, Fig 1A).

Genetic architecture of ischemic stroke in Brazilian Sickle Cell

We observed 28 genome-wide significant variants (Supplemental Table 1) across 14 independent regions associated with stroke occurring at younger ages (Figure 1B, Table 2). No systemic inflation of *P*-values was observed (Genomic Control Lambda = 1.02; Supplemental Figure 2). Most variants sites were intergenic; however, three regions were intronic to genes *CUX1*, *KIAA1217*, and *CNNM2* (Supplemental Figure 3). Also Included in this list of 14 lead SNPs were three regions previously linked to stroke – *ADAMTS2*, *CDK18*, *FUT8*^{18–20}.

Out of the 14 genome-wide lead variants, 5 were only observed in heterozygous state whereas 9 were observed in both heterozygous and homozygous variant state. Individuals

possessing at least one of these 14 variants exhibited increased risk for ischemic stroke at earlier ages (P < 0.001 for all 14 variants, Gehan-Wilcoxon) (Supplemental Figure 4).

Multiple variant survival analysis

To investigate the impact of possessing multiple lead risk variants within the same individual, participants were divided into categories based on the number of lead SNPs they harbored. Individuals with zero lead SNPs had a stroke rate of 4% (n=716), whereas those with at least one of the 14 lead variants had an increased rate at 14% (n=430), and this rate increased even further for individuals with two (36%; n=125), three (70%; n=50), or four or more variants (83%; n=12) (Figure 2A). The relative impact of these alleles on cohort level risk was measured by calculating the product of minor allele frequency with the hazard ratio (Figure 2B). The variant rs12144136 near *CDK18* exhibited the highest relative weight of risk for early stroke within this cohort.

Cis-co-localization analysis of ischemic stroke using GTEx

All 14 candidate SNPs from this study occur in inter-genic or intronic regions. To investigate evidence of these SNPs influencing gene expression, the lead SNPs were searched against the GTEx database of gene expression to identify *cis-* and *trans-* expression quantitative trail loci (eQTL) variant associations across different tissue types. The lead SNP on chromosome 1, rs12144136, is a *cis-*eQTL for the gene *CDK18* within the Skeletal Muscle tissue category (Supplemental Table 2) which appears to upregulate this gene. No other SNPs at genome-wide significance showed evidence of *cis-*eQTL activity.

Expanding to 1Mb regions around each lead SNP, we assessed co-localization of *cis*-eQTL associations with 24 different tissue types relevant to stroke. Three of the 14 candidate regions showed strong evidence (Posterior Probability, PP > 0.9) of colocalization with a *cis*-eQTL (Supplemental Table 2). On chromosome 12, two *cis*-eQTLs were found for genes *LUM* in the frontal cortex (rs77217583) and *NUDT4* (rs117881990) in the hypothalamus. On chromosome 14, one *cis*-eQTL was found for *RAB15* in the substantia nigra; and on chromosome 15, one *cis*-eQTL was found for *VPS13C* in cervical spinal cord. Three other moderately strong signals (P > 0.5) were detected on chromosome 8 – one *cis*-eQTL for *WRN* expression in heart left ventricle and two *cis*-eQTLs for *PURG* in the substantia nigra.

Gene-level Association

We performed a gene-level association on 19,021 protein coding genes using the 14M site summary statistics (Supplemental Figure 5; Supplementary Table 3). No genes exhibited genome-wide significant enrichment defined at the Bonferroni-corrected alpha of 2.6×10^{-6} ; however, three genes exhibited some evidence of association ($P < 10^{-4}$), including *GPR15* (P= 3.9×10^{-5}), *MMP26* (P= 4.6×10^{-5}), and *GOLT1B* (P= 7.5×10^{-5}).

Gene-set enrichment analysis was performed to predict impact on biological functions using the Gene Ontology (GO) database (Supplemental Table 4). While no gene set achieved the Bonferroni-corrected p-value of 5×10^{-6} , top results at the *P*<0.001 threshold included the platelet dense tubular network (*P*= 3.63×10^{-5}), the hemoglobin complex (*P*=0.0007), gas

transport (P = 0.0006), and regulation of systemic arterial blood pressure by hormone (P = 0.0008).

Replication with other SCD cohorts

Candidate SNP analysis of the 14 lead SNPs was performed within two independent SCD cohorts: Walk-PHaSST (12 cases, 300 controls)³⁵ and PUSH (14 cases, 202 controls)³⁶ (Supplemental Table 5). Cox regression for stroke timing, adjusting for sex and the first genetic principal component, showed increased risk for early stroke at SNPs rs1209987 near gene *TBC1D32* (P= 0.01) and rs188599171 near gene *CUX1* (P= 0.001) within the combined Walk-PHaSST/PUSH cohort. Meta-analysis with the REDS-III Brazil SCD cohort and the combined Walk-PHaSST/PUSH cohort showed genome-wide significance for four SNPs: rs1209987 near gene *TBC1D32* (P = 3.36×10–10), rs188599171 near *CUX1* (P = 5.89×10–11), rs77900855 near *BTG1* (P=4.66×10–8), and rs141674494 near *VPS13C* (1.68×10⁻⁹), although we note these results are being driven largely by the REDS-III Brazil SCD cohort as none of the SNPs reached genome-wide significance within Walk-PHaSST or PUSH.

A comparison was also made to the results from Flanagan et al¹³ (122 cases, 167 controls). None of the 14 lead SNPs were genotyped in this cohort, and investigating the surrounding 1Mb regions did not uncover any loci exhibiting association with ischemic above the P=0.001 threshold (results not shown).

A third and final comparison was made to an independent cohort of SCD children with genome sequence data, comprised of 15 patients with a history of ischemic stroke, 26 patients with a history of abnormal TCD, and 140 control patients (J.M.F. and R.E.W. Unpublished data, 2022). A combined Armitage test for trend on the combined stoke/ abnormal TDC (N=413) versus controls (N=140) showed modest, but not genome-wide significant, replication of rs116211928 near gene *TLE6* (P= 0.022).

Candidate-gene Enrichment Analysis to GWAS Catalog

To assess overlap in genetic architecture between early ischemic stroke and other relevant cardiovascular traits, we performed an enrichment test against the GWAS Catalog of gene by trait associations. Candidate genes from the current study were defined as located within 500Kb of a lead SNP, or the most proximal gene if none were present in that region, resulting in a list of 52 genes (Supplemental Table 6). Significant overlap in gene sets was observed for mean arterial pressure (3/9 genes; $P = 5.82 \times 10^{-4}$), hypertension (5/98 genes; $P = 1.48 \times 10^{-3}$), immature fraction of reticulocytes (4/104 genes; $P = 1.64 \times 10^{-2}$), and myocardial infarction (3/55 genes; $P = 3.53 \times 10^{-2}$) (Figure 3).

Comparison to previous ischemic stroke studies in SCD and non-SCD

Previous candidate SNP studies of stroke in SCD have evaluated 12 SNPs, as well as one additional SNP from a GWAS^{8–15}. None of these candidates SNPs reached genome-wide significance in the current study (Supplemental Table 7). Expanding the search to surrounding 1Mb region centered on the 13 candidate SNPs uncovered the lead SNP rs147625068 ($P = 3.7 \times 10^{-9}$) near gene *LTC4S* on chromosome 9.

A similar search was performed for non-SCD early onset ischemic stroke candidate genes: *ADAMTS2* (rs469568), *ADAMTS12* (rs1364044), *CDK18* (rs77571454), *HABP2* (rs11196288)^{17–20}. In the current study, the variant rs77571454 near gene *CDK18* was not observed, and none of the other three variants reached genome-wide significance (Supplemental Table 8). Searching within the 1Mb region centered on these four candidate SNPs uncovered the two lead SNPs rs12144136 ($P = 2.38 \times 10^{-9}$), near *CDK18*, and rs147625068 ($P = 3.7 \times 10^{-9}$), near *ADAMTS2*, and one additional variant near *ADAMTS12* (rs534089428, $P = 2.43 \times 10^{-5}$).

DISCUSSSION

We have discovered 14 genetic regions associated with ischemic stroke occurring at earlier ages in SCD individuals. This study included 1,333 *HbSS* individuals, the largest genome-wide evaluation of ischemic stroke within SCD to date. We implemented a Cox proportional hazards model to estimate risk at earlier timepoints. Results implicate gene candidates previously linked with non-SCD early-onset stroke – *ADAMTS2*, *CDK18*, genes linked to cardiovascular disease risk factors – *BTG1*, *CNMM2*, *CUX1*, *FUT8*, *KIAA1217*, *LUM*, *NT5C2*, *NUDT4*, *PKD1L1*, *WRN*; and eight novel genes – *SH3RF3*, *RAB15*, *RPS3AP52*, *TBC1D32*, *VPS13C*, and *TLE6*. To our knowledge this is the first independent replication of genetic risk associated with increased stroke at alleles near genes *CDK18* and *FUT8*.

Previous candidate gene studies of stroke in SCD implicated five alleles within genes *TGFBR3, TEK, ANXA2, IL4R,* and *ADCY9*^{10–12}; however, the single published GWAS of ischemic stroke in SCD did not find evidence for alleles near these genes influencing stroke risk but instead found alleles near two novel genes – *GOLGB1* and *ENPP1*¹³. We did not find evidence of alleles near these genes exhibiting genome-wide significant association to stroke. The reasons for this are unclear, although we should note the previous GWAS enrolled African Americans whereas the current work enrolled Brazilians, each having distinct histories of admixture.

Results from this work implicate genes previously linked to non-SCD early onset stroke: *ADAMTS2* and *CDK18*^{17–19}. Alleles near genes *ADAMTS2* and *ADAMTS12* were previously associated with pediatric stroke in two separate U.S. populations^{17,18}. *ADAMTS2* also appears to have a role in cerebral aneurysm³⁷ and cardiac muscle hypertrophy³⁸ possibly through regulation of the extracellular matrix³⁹. We observed one variant, rs534089428, within 500kb of *ADAMTS12* at $P = 2 \times 10^{-5}$.

High reticulocyte count is a reliable risk factor for ischemic stroke in sickle cell individuals¹. Variants affecting candidate genes from the current study, *CUX1, CNNM2*, and *NUDT4*, have been previously associated with higher reticulocyte counts^{40 41}. *CNNM2* has also been linked to numerous stroke related traits – hypertension, coronary artery disease, and mean corpuscular hemoglobin^{42–49}. *CNNM2* gene is expressed throughout the body but is particularly high in the brain, kidney, and endocrine tissue⁵⁰ and appears to play a role in renal magnesium uptake^{51,52}

This study is limited in a few ways. First, this work did not consider the influence of environmental exposures and lifestyles on ischemic stroke risk. Non-acute, unreported comorbidities may also play a role. This study is larger than previous at N=1,333, but the observed association of alleles at a frequency of 1-5% within a case size of N=178 means that relatively few individuals are driving this signal, and future studies in independent cohorts will need to validate these findings. In addition, we define candidate genes based on proximity to lead SNPs and note the possibility risk alleles may actually be linked to distant genes via, for example, trans-QTL effects.

In conclusion, we have identified risk alleles associated with earlier ischemic stroke. Findings from this work have overlapped previously identified non-SCD early onset ischemic stroke genes – ADAMTS2 and CDK18– and discovered 16 novel genes. Future studies will be needed to confirm the clinical significance of these findings and to disentangle the potentially complex interaction among these alleles.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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EJE, SK, GPP wrote the paper; GPP, CD, BC, SK, ES designed the study; EJE, FF, GPP performed the analysis; CSA, DOWR, DTSC collected data; JMF, REW, XZ, VG, MG, YZ, MN, SN provided lookup analysis for independent SCD cohorts. Special thanks to patients and research staff at each hemocenter in Brazil participating in this study. We recognize the following people for their commitment and contribution to this project: Alfredo Mendrone Jr., Cesar de Almeida Neto, Roberta Calcucci, Erivanda Bezerra, Carolina Miranda, Franciane Mendes de Oliveira, Valquiria Reis, Nayara Durte, Barbara Malta, Jose Wilson Sales, Maria Aparecida Souza, Rodrigo Ferreira, Maria do Carmo Valgueir, Regina Gomes, Airly Goes Maciel, Rebeca Talamatu Dantas, Flavia Herculano, Ana Claudia Pereira, Ana Carla Alvarenga, Adriana Grilo, Fabiana Canedo, Pedro Losco Takecian, Mina Cintho Ozahata, Rodrigo Muller de Carvalho, Christopher McClure, Simone A. Glynn.

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Figure 1.

(A) Histogram of age at first ischemic stroke for the GWAS cohort. (B) Manhattan plot of results from proportional hazards GWAS P-values for ischemic stroke in HbSS SCD (N=1,333). Horizontal dotted line represents the genome-wide significance threshold of $P = 5 \times 10$ –8. The nearest gene to each lead genome-wide significant SNP is annotated.



Figure 2.

(A) Kaplan-Meier plot of individuals harboring multiple co-occurring genome-wide significant risk alleles for early stroke. Curves represent subsets of the study cohort who possessed zero (N=716), one (N=430), two (N=125), three (N=50), or four or more (N=12) stroke risk alleles in any combination from the 14 lead SNPS. (+) represents right-censored data. (B) A scatter plot of observed hazard ratios (HR) and minor allele frequencies (MAF). Points represent the 14 lead SNPs and are annotated with the nearest gene. The relative impact of each allele on the cohort-scale risk of earlier ischemic stroke is calculated as the product of MAF and HR and is color coded from gray (lower impact) to black (higher impact).



Figure 3.

Gene enrichment test results against the GWAS Catalog of reported gene by trait associations. Candidate genes were defined as being within 500kb of a lead SNP. Left, horizontal bar plots of the proportion of genes within each trait category observed in the candidate gene list (gray) and the P values (black) from a hypergeometric enrichment test (FDR adjusted). Traits relevant to ischemic stroke are listed in bold. Right, a visual matrix showing which candidate genes overlap a given trait.

Table 1.

Demographics of HbSS Individuals within the REDS-III Brazil SCD Cohort

Total cohort size	N=1,333
Sex (%)	
Female	720 (54.0)
Male	613 (46.0)
Age at enrollment, mean (\pm SD), y	20.6 (±13.1)
Age at first stroke, mean (±SD), y	9.5 (±9.4)
Stroke, no. (%)	
cases	178 (13.4)
controls	1155 (86.6)
Hemocenter, no. (%)	
Hemominas BH	313 (23.5)
Hemominas JFO	149 (11.2)
Hemominas MOC	205 (15.4)
Hemope	285 (21.4)
Hemorio	332 (24.9)
ITACI SP	49 (0.04)

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Chr	Position (b38)	Effect Allele	Nearest Protein Coding Gene	rsID	P-Value	Effect AF (%)	HR	95% CI	Variant Type	Functional Annotation
1	205,490,782	G	CDK18	rs12144136	$2.38 imes 10^{-9}$	5.1	2.7	2.4 - 3.0	SNP	Intergenic
2	109,059,503	А	SH3RF3	rs10181988	3.35×10^{-8}	2.7	3.2	2.8 - 3.6	SNP	Intergenic
5	179,375,279	А	ADAMTS2	rs147625068	$3.70 imes 10^{-9}$	1.0	5.4	4.8 - 5.9	SNP	Intergenic
9	120,572,043	Т	TBC1D32	rs1209987	2.52×10^{-9}	10.3	2.3	2.0 - 2.6	SNP	Intergenic
٢	47,681,969	С	PKDILI	rs181930335	$2.71 imes 10^{-8}$	2.3	3.6	3.2 - 4.1	SNP	Intergenic
٢	102,180,550	А	CUXI	rs188599171	$3.70 imes 10^{-9}$	1.1	5.6	5.0 - 6.2	SNP	Intronic
8	31,319,867	А	WRN	rs115684945	$3.10 imes 10^{-8}$	1.8	3.4	3.0 - 3.8	SNP	Intergenic
10	23,744,565	А	<i>KIAA1217</i>	rs115292858	4.43×10^{-8}	2.2	3.8	3.3 - 4.3	SNP	Intronic
10	102,993,950		CNNM2, NT5C2	rs367825668	$3.90 imes 10^{-8}$	1.7	3.4	3.0 - 3.9	DEL	Intronic
12	92,093,367	Ū	BTGI	rs77900855	$2.51 imes 10^{-8}$	1.0	5.0	4.5 - 5.6	SNP	Intergenic
13	68,609,784	С	RPS3AP52	rs73204086	3.78×10^{-12}	1.1	5.5	5.0 - 6.0	SNP	Intergenic
14	65,317,148	Н	FUT8	rs150572699	$2.58 imes 10^{-9}$	1.4	3.9	2.5 - 6.2	SNP	Intergenic
15	61,683,898	G	VPS13C	rs141674494	5.40×10^{-10}	1.1	6.3	5.7 - 6.9	SNP	Intergenic
19	2,961,825	Ū	TLE6	rs116211928	2.25×10^{-8}	1.4	4.8	4.2 - 5.3	SNP	Intergenic
5										

Confidence Interval; Genomic positions in GRCh38 build. 80% 52% CI = Hazard Katio; Chr = Chromosome; AF = Allele Frequency; HK