Sickle cell trait and risk of cognitive impairment in African-Americans: The REGARDS cohort

Christina R. Cahill, University of Vermont
Justin M. Leach, University of Alabama at Birmingham
Leslie A. McClure, Drexel University
Marguerite Ryan Irvin, University of Alabama at Birmingham
Neil A. Zakai, University of Vermont
Rakhi Naik, Johns Hopkins University
Frederick Unverzagt, Indiana University
Virginia G. Wadley, University of Alabama at Birmingham
Hyacinth Hyacinth, Emory University
Jennifer Manly, Columbia University Medical Center

Only first 10 authors above; see publication for full author list.

Journal Title: EClinicalMedicine
Volume: Volume 11
Publisher: Elsevier | 2019-05-01, Pages 27-33
Type of Work: Article | Final Publisher PDF
Publisher DOI: 10.1016/j.eclinm.2019.05.003
Permanent URL: https://pid.emory.edu/ark:/25593/trwkc

Final published version: http://dx.doi.org/10.1016/j.eclinm.2019.05.003

Copyright information:
© 2019
This is an Open Access work distributed under the terms of the Creative Commons Attribution-NonCommercial-NoDerivatives 4.0 International License (http://creativecommons.org/licenses/by-nc-nd/4.0/).

Accessed April 5, 2022 9:43 PM EDT
Sickle cell trait and risk of cognitive impairment in African-Americans: The REGARDS cohort

Christina R. Cahill a, Justin M. Leach b, Leslie A. McClure c, Marguerite Ryan Irvin b, Neil A. Zakai d,e, Rakhi Naik f, Frederick Unverzagt g, Virginia G. Wadley h, Hyacinth I. Hyacinth i, Jennifer Manly j, Suzanne E. Judd b, Cheryl Winkler k, Mary Cushman d,e,*

a Larner College of Medicine at the University of Vermont, Burlington, VT, United States of America
b Department of Epidemiology, University of Alabama at Birmingham, Birmingham, AL, United States of America
c Department of Biostatistics, University of Alabama at Birmingham, Birmingham, AL, United States of America
d Department of Biostatistics, Dornsife School of Public Health, Drexel University, Philadelphia, PA, United States of America
e Department of Medicine, Larner College of Medicine at the University of Vermont, Burlington, VT, United States of America
f Department of Medicine, Johns Hopkins University, Baltimore, MD, United States of America
g Department of Pathology and Laboratory Medicine, Larner College of Medicine at the University of Vermont, Burlington, VT, United States of America
h Department of Psychiatry, Indiana University School of Medicine, Indianapolis, IN, United States of America
i Department of Medicine, Division of Gerontology Geriatrics and Palliative Care, University of Alabama at Birmingham, Birmingham, AL, United States of America
j The Taub Institute for Research in Alzheimer’s Disease and the Aging Brain, Columbia University Medical Center, New York, NY, United States of America
k Molecular Genetics Epidemiology Section, Frederick National Laboratory for Cancer Research, National Cancer Institute, National Institutes of Health, Frederick, MD, United States of America

A R T I C L E   I N F O

Article history:
Received 14 October 2018
Received in revised form 2 May 2019
Accepted 3 May 2019
Available online 24 May 2019

Keywords:
Sickle cell trait
Cognitive dysfunction
Cognition
Prospective studies
Risk factors
Epidemiology

A B S T R A C T

Background: Sickle cell anemia may be associated with cognitive dysfunction, and some complications of sickle cell anemia might affect those with sickle cell trait (SCT), so we hypothesized that SCT is a risk factor for cognitive impairment.

Methods: The Reasons for Geographic and Racial Differences in Stroke (REGARDS) study enrolled a national cohort of 30,239 white and black Americans from 2003 to 7, who are followed every 6 months. Baseline and annual global cognitive function testing used the Six-Item Screener (SIS), a validated instrument (scores range 0–6; ≤4 indicates cognitive impairment). Participants with baseline cognitive impairment and whites were excluded. Logistic regression was used to calculate the association of SCT with incident cognitive impairment, adjusted for risk factors. Linear mixed models assessed multivariable-adjusted change in test scores on a biennially administered 3-test battery measuring learning, memory, and semantic and phonemic fluency.

Findings: Among 7743 participants followed for a median of 7·1 years, 85 of 583 participants with SCT (14·6%) developed incident cognitive impairment compared to 902 of 7160 (12·6%) without SCT. In univariate analysis, the odds ratio (OR) of incident cognitive impairment was 1·18 (95% CI: 0·93, 1·51) for those with SCT vs. those without. Adjustment did not impact the OR. There was no difference in change on 3-test battery scores by SCT status (all p > 0·11).

Interpretation: In this prospective cohort study of black Americans, SCT was not associated with incident cognitive impairment or decline in test scores of learning, memory and executive function.

Funding: National Institutes of Health, American Society of Hematology.

© 2019 Published by Elsevier Ltd. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).

1. Introduction

Vascular risk factors are also risk factors for cognitive decline and impairment, mediated partly by small, subclinical strokes [1–4]. Patients with sickle cell anemia (SCA) may develop impaired cognitive function. Specifically, children [5] and adults [6] with SCA score lower on cognitive test scores than controls and children with SCA are at risk for lower academic attainment [7]. Silent cerebral infarction is common in children and adults with SCA [8], and white matter hyperintensities, a measure of silent cerebral infarction, correlate with poorer neurocognitive outcomes in these children [9]. SCA has also been linked with cognitive processing speed independent of silent infarcts, but related to MRI-defined white matter integrity [10]. In a study of adults with SCA cortical and subcortical brain volumes were lower than controls, and these lower volumes correlated with lower cognitive performance [11].

DOI of original article: https://doi.org/10.1016/j.eclinm.2019.04.006.

* Corresponding author: Department of Medicine, Larner College of Medicine at the University of Vermont, 360 South Park Drive, Colchester, VT 05446, United States of America.
E-mail address: mary.cushman@uvm.edu (M. Cushman).
2. Methods

2.1. Study participants and data collection

The REGARDS study is a longitudinal observational study investigating racial and geographic variation in incidence of stroke and acquired cognitive impairment in the contiguous United States. Details of the study design were published elsewhere [26]. Cohort participants were randomly selected by mail and enrolled by telephone followed by an in-home visit between 2003 and 7. The aim was to enroll 50% non-Hispanic black and 50% non-Hispanic white participants aged 45 and older, with 50% residing in the southeast. Exclusion criteria included individuals who indicated race other than black or white, cognitive impairment precluding ability to complete a telephone interview, active cancer within 1 year or undergoing treatment for cancer, a medical condition preventing long term follow-up, residing in or waiting for nursing home residence, or inability to communicate in English [26]. Enrollment results yielded a cohort with 51% female and 42% black participants [26].

Baseline participant characteristics were obtained via a computer-assisted telephone interview followed by in-home examination using a standard protocol that included phlebotomy and shipment of blood and urine samples to a central laboratory for storage and measurement of glucose, lipid profile and kidney function [27]. SCT was determined via genotyping using a TaqMan SNP Genotyping Assay (Applied Biosystems/ThermoFisher Scientific) [16]. Among a subset of participants with available genomic data, ten principal components of ancestry were determined using EIGENSTRAT to control for population stratification [17,28].

The study methods were approved by the institutional review boards at all participating institutions and all participants provided informed consent. Boards included the University of Alabama at Birmingham Institutional Review Board for Human Use, the University of Vermont Research Protections Office, the University of Cincinnati Human Research Protection Program, the Wake Forest University Institutional Review Board, and the Columbia University Human Research Protection Office.

2.2. Covariate measurements

Race was determined by participant self-report as black or white. Age, sex, education, household income level, and region of residence were determined by self-report. Education was categorized as less than high school, high school graduate, some college, or college graduate and above. Income was categorized as $20,000/yr, $20,000 to $34,999/yr, $35,000 to $74,999/yr, or >$75,000/yr. Current cigarette smoking, alcohol use, exercise frequency, and use of medications were determined by interview. Hypertension was defined as baseline systolic blood pressure ≥ 140 mmHg, diastolic blood pressure ≥ 90 mmHg or self-reported use of antihypertensive medications. Hyperlipidaemia was defined as low-density lipoprotein > 130 mg/dl, or self-reported use of a cholesterol-lowering medication. Diabetes was defined by a fasting glucose > 126 mg/dl, nonfasting glucose > 200 mg/dl, or self-reported use of antidiabetes medications. Coronary heart disease was determined by self-reported myocardial infarction (MI), coronary artery bypass graft, angioplasty or stenting, or evidence of MI from baseline electrocardiogram. Atrial fibrillation was defined as self-reported or via electrocardiogram evidence. Left ventricular hypertrophy (LVH) was defined by electrocardiogram. Estimated glomerular filtration rate (eGFR) was calculated from serum creatinine using the Chronic Kidney Disease Epidemiology Collaboration equation [29].

Table 1. Flow diagram of participant inclusion.

| Total REGARDS Participants: 30 239 |
|-----------------------------------|
| Excluded white participants: 17 725 |
| Black participants: 12 514 |
| Excluded those with missing SCT genotyping (1 935), hemoglobin S5 (5) or SC (1) and with missing data on baseline cognitive impairment or having only one assessment (874) |
| 9 699 black participants |
| Participants included in analysis: 7 743 (583 with SCT) |
| Excluded those with baseline cognitive impairment (1 338) and with incident stroke (618) |

Fig. 1. Flow diagram of participant inclusion.
2.3. Cognitive function assessments

Cognitive outcomes were studied in all participants in two ways, considered here as co-primary outcomes: incident cognitive impairment on a test of global cognitive function and longitudinal change of cognitive domain test scores reflecting learning, memory and executive function.

The study conducted global cognitive function testing using the Six-item Screener (SIS), a validated telephone-administered instrument for global cognitive function that assesses 3-item recall and orientation to year, month, and day of the week, yielding a score from 0 to 6 correctly answered questions [30,31]. A score ≤4 is considered positive for cognitive impairment. The SIS was administered at baseline and then annually to all participants. The outcome of incident cognitive impairment was defined at the most recent assessment as of April 1, 2015, when our analysis data set was closed.

Participants also completed a 3-test battery every two years that evaluated measures of learning, memory and executive function [27]. Validated instruments were telephone-administered, including the Consortium to Establish a Registry for Alzheimer’s Disease (CERAD) battery to assess word list learning (WLL), delayed recall by word list recall (WLR), semantic fluency by the Animal Fluency Test (AFT) score, and phonemic fluency by the Letter F test [32–34]. WLL is the sum of words learned over 3 trials, where participants are tested on immediate recall of 10 words, with scores ranging from 0 to 30. WLR is the sum of the same words recalled after a delay filled with intervening questions, with a range of 0–10 words. The AFT score is determined by the number of animals a participant can name within 1-min. The Letter F score is

| Covariate (N missing) | All participants (N = 7743) | SCT (N = 583) | No SCT (N = 7160) | p-Value |
|-----------------------|-----------------------------|---------------|-------------------|---------|
| Age, years (0)        | 63·1 (8·9)                  | 62·9 (9·0)    | 63·1 (8·9)        | 0·48    |
| eGFR, ml/min/1·73 m²  | 90 (22)                     | 86 (24)       | 90 (22)           | <0·0001 |
| Systolic Blood Pressure, mm Hg (21) | 130 (17)       | 131 (17)      | 130 (17)          | 0·33    |
| Log(ACR), mg/g (282)  | 2·4 (1·1)                   | 2·7 (1·4)     | 2·4 (1·3)         | <0·0001 |
| Sex (0)               | Female 4898 (63·0%)         | 388 (66·6%)   | 4510 (63·0%)      | 0·09    |
|                       | Male 2845 (37·0%)           | 195 (33·4%)   | 2650 (37·0%)      |        |
| Education (5)         | High School 1225 (15·8%)   | 90 (15·4%)    | 1135 (15·9%)      | 0·98    |
|                       | High School graduate 2110 (27·3%) | 158 (27·3%) | 1952 (27·3%)      |        |
|                       | Some College 2165 (28·0%)  | 167 (28·6%)   | 1998 (28·7%)      |        |
|                       | College graduate 2238 (28·9%) | 168 (28·8%)  | 2070 (28·9%)      |        |
| Income (0)            | <$20 k 1814 (23·4%)         | 153 (26·2%)   | 1661 (23·2%)      | 0·26    |
|                       | $20 k-$34 k 2018 (26·1%)    | 145 (24·9%)   | 1873 (26·2%)      |        |
|                       | $35 k-$74 k 2213 (28·6%)    | 150 (25·7%)   | 2063 (28·6%)      |        |
|                       | >$75 k 833 (10·8%)          | 62 (10·6%)    | 771 (10·8%)       |        |
| Smoking status (36)   | Current 1283 (16·6%)        | 91 (15·7%)    | 1192 (16·7%)      | 0·03    |
|                       | Past 2833 (36·8%)           | 189 (32·5%)   | 2644 (37·1%)      |        |
|                       | Never 3591 (46·6%)          | 301 (51·8%)   | 3290 (46·2%)      |        |
| Exercise (93)         | 1-3 Times/Week 2934 (38·4%) | 221 (38·2%)   | 2713 (38·4%)      |        |
|                       | 4+ Times/Week 2018 (26·4%)  | 159 (27·5%)   | 1859 (26·3%)      |        |
| Coronary heart disease (135) | No 6616 (87·0%) | 495 (66·7%) | 6121 (87·0%) | 0·84 |
|                       | Yes 992 (13·0%)             | 76 (13·3%)    | 916 (13·0%)       |        |
| Left ventricular hypertrophy (131) | No 6582 (86·3%) | 484 (84·6%) | 6098 (86·5%) | 0·22 |
|                       | Yes 1, 044 (13·7%)          | 88 (15·4%)    | 956 (13·5%)       |        |
| Diabetes (78)         | No 5563 (72·6%)             | 409 (71·1%)   | 5154 (72·7%)      | 0·42    |
|                       | Yes 2102 (27·4%)            | 166 (28·9%)   | 1936 (27·3%)      |        |
| Atrial fibrillation (192) | No 7001 (92·7%) | 518 (91·4%) | 6483 (92·6%) | 0·20 |
|                       | Yes 550 (7·3%)              | 49 (8·6%)     | 501 (7·2%)        |        |
| Hyperlipidaemia (73)  | No 5366 (70·0%)             | 398 (68·6%)   | 4968 (70·1)       | 0·46    |
|                       | Yes 2304 (30·0%)            | 182 (31·4%)   | 2122 (29·9%)      |        |
| Statin Use (25)       | No 5576 (72·2%)             | 411 (70·5%)   | 5165 (72·4%)      | 0·33    |
|                       | Yes 2142 (27·8%)            | 172 (29·5%)   | 1970 (27·6%)      |        |
| Hypertension (289)    | No 2700 (36%)               | 214 (38·7%)   | 2486 (36·0%)      | 0·21    |
|                       | Yes 4754 (64%)              | 339 (61·3%)   | 4415 (64·0%)      |        |
determined by the number of words beginning with the letter F that a participant can say within 1-min.

### 2.4. Inclusion/Exclusion criteria for analysis

We included black participants who had available data on baseline and at least one follow up SIS, and with SCT genotyping. Participants with hemoglobin SS or SC genotype, baseline cognitive impairment (SIS ≤ 4), or who developed incident stroke were excluded from all analyses.

#### Table 2

Baseline characteristics by incident cognitive impairment on the six-item screener.

| Covariate (N missing) | All participants (N = 7743) | Incident impairment (N = 987) | No incident impairment (N = 6756) |
|-----------------------|-----------------------------|-------------------------------|----------------------------------|
| Age, years (0)        | 63.1 (8.9)                  | 68.6 (9.2)                   | 62.3 (8.6)                       |
| eGFR, ml/min/1·73 m² (48) | 90 (22)                    | 82 (23)                      | 91 (22)                          |
| Systolic Blood Pressure, mm Hg (21) | 130 (17)                  | 133 (18)                     | 130 (17)                         |
| Log(ACR), mg/g (282) | 2.40 (1.3)                  | 2.64 (1.5)                   | 2.34 (1.3)                       |
| Sex (0)               | Female 4898 (63.0%)         | 541 (54.8%)                  | 4357 (64.5%)                     |
|                       | Male 2845 (37.0%)           | 446 (45.2%)                  | 2399 (35.5%)                     |
| Education (5)         | High School 1225 (15.8%)   | 271 (27.5%)                  | 954 (14.3%)                      |
|                       | Some College 2165 (28.0%)  | 236 (24.0%)                  | 1929 (28.6%)                     |
|                       | College graduate 2238 (28.9%) | 190 (19.3%)                 | 2048 (30.3%)                     |
| Income (0)            | <$20k 1814 (23.4%)          | 335 (33.9%)                  | 1479 (21.9)                      |
|                       | $20k–$34k 2018 (26.1%)     | 283 (28.7%)                  | 1735 (25.7%)                     |
|                       | $35k–$74k 2213 (28.6%)     | 179 (18.1%)                  | 2034 (30.1%)                     |
|                       | $75k+ 833 (10.8%)           | 49 (5.0%)                    | 784 (11.6%)                      |
| Refused (0)           | 865 (11.2%)                 | 141 (14.3%)                  | 724 (10.7%)                      |
| Region (0)            | Belt 2565 (33.1%)           | 323 (32.7%)                  | 2242 (33.2%)                     |
|                       | Buckle 1359 (17.6%)         | 163 (15.6%)                  | 1196 (17.7%)                     |
|                       | NonBelt 3819 (49.3%)        | 501 (50.8%)                  | 3318 (49.1%)                     |
| Alcohol use group (199) | Heavy 189 (2.5%)           | 26 (2.7%)                    | 163 (2.5%)                       |
|                       | Moderate 2047 (27.1%)       | 213 (22.2%)                  | 1834 (27.9%)                     |
|                       | None 5308 (70.4%)           | 721 (75.1%)                  | 4587 (69.7%)                     |
| Smoking status (36)   | Current 1283 (16.6%)        | 157 (16.0%)                  | 1126 (16.7%)                     |
|                       | Past 2833 (36.3%)           | 370 (37.7%)                  | 2463 (36.6%)                     |
|                       | Never 3591 (46.6%)          | 455 (46.3%)                  | 3136 (46.6%)                     |
| Exercise (93)         | Physical activity 3 Times/Week 2934 (38.4%) | 346 (35.6%) | 2588 (38.7%) |
|                       | None 2698 (35.3%)           | 367 (37.8%)                  | 2331 (34.9%)                     |
| Coronary heart disease (135) | No 6616 (87.0%)        | 785 (80.8%)                  | 5831 (87.9%)                     |
|                       | Yes 992 (13.0%)             | 187 (19.2%)                  | 805 (12.1%)                      |
| Left ventricular hypertrophy (131) | No 6582 (86.3%)        | 810 (83.3%)                  | 5772 (86.8%)                     |
|                       | Yes 1,044 (13.7%)           | 163 (16.8%)                  | 881 (13.2%)                      |
| Diabetes (78)         | No 5563 (72.6%)             | 661 (67.4%)                  | 4902 (73.3%)                     |
|                       | Yes 2102 (27.4%)            | 320 (32.6%)                  | 1782 (26.7%)                     |
| Atrial fibrillation (192) | No 7001 (92.7%)        | 885 (94.2%)                  | 6116 (92.8%)                     |
|                       | Yes 550 (7.3%)              | 73 (7.6%)                    | 477 (7.2%)                       |
| Hyperlipidaemia (73)  | No 5366 (70.0%)             | 632 (64.6%)                  | 4734 (70.7%)                     |
|                       | Yes 2304 (30.0%)            | 346 (34.4%)                  | 1958 (29.3%)                     |
| Statin (25)           | No 5576 (72.2%)             | 677 (76.8%)                  | 4899 (78.8%)                     |
|                       | Yes 2142 (27.8%)            | 308 (33.2%)                  | 1834 (27.2%)                     |
| Hypertension (289)    | No 2700 (36.3%)             | 306 (32.7%)                  | 2394 (36.7%)                     |
|                       | Yes 4754 (64%)              | 629 (67.3%)                  | 4125 (63.3%)                     |

#### 2.5. Statistical analysis

Differences in baseline characteristics by SCT and incident cognitive impairment on the SIS were analyzed by t-tests for continuous measures and chi-square tests for categorical measures. For incident cognitive impairment based on the SIS, we used logistic regression to calculate odds ratios (OR) with 95% confidence intervals (CI) by SCT status. Statistical significance was defined as p ≤ 0.05. Linear mixed models were used to study association of WLL with WLR and semantic and verbal fluency over time. Similar to prior REGARDS reports, no random effects accounting for time between tests were included [35]. For both types of analysis, multivariable models were fitted to adjust for age, sex, education, income, region of residence, eGFR, systolic blood pressure, alcohol use, smoking status, exercise frequency, coronary heart disease, LVH, atrial fibrillation, hyperlipidaemia, statin use, diabetes, and hypertension. Models were repeated following removal of variables that were not significantly associated with the cognitive outcome. Censoring occurred at death or withdrawal from the study.

The first sensitivity analysis addressed whether observed associations might be explained by other genetic factors associated with African ancestry that are in linkage with SCT. Specifically, this was done on the subset with available genetic ancestry information by adding adjustment for African ancestry using the first ten principal components of ancestry. A second sensitivity analysis for the association of cognitive impairment by the SIS only, evaluated the cross-sectional association of SCT with a SIS ≤ 4 compared to 5 or 6 using analogous regression models to those for incident cognitive impairment in 9549 participants. This analysis involved baseline SIS scores, but included participants whose baseline impairment or lack of follow up assessments excluded them from the analysis of incident impairment.

Interactions between SCT and age (continuous variable), sex, and diabetes status were tested for in all analyses. Analysis was performed with SAS 9.4.

---

Fig. 2. Sickle cell trait and longitudinal change in Word List Learning (WLL) and Delayed Recall (WLR). Blue lines represent those with sickle cell trait and black lines those without sickle cell trait. Models were adjusted for age, sex, education, income, region, eGFR, systolic blood pressure, diabetes, alcohol use, smoking status, exercise frequency, coronary heart disease, left ventricular hypertrophy, atrial fibrillation, hyperlipidaemia, statin use, and hypertension. Note, graph starts with a score of 5 words for ease of displaying the data. Numbers below the figure show the sample sizes. Baseline SD of Scores: WLL (SCT) = 5.0; WLL (No SCT) = 5.0; WLR (SCT) = 2.3; WLR (No SCT) = 2.2. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
2.6. Role of the funding source

The funding source had no role in the study design, collection, analysis and interpretation of data, authorship or decision to submit for publication.

2.7. Data statement

There are restrictions to the open publication of a REGARDS dataset but data may be requested per study policies at www.regardsstudy.org.

3. Results

Fig. 1 shows a flow diagram of participant inclusion in the analysis of incident cognitive impairment by the SIS. Baseline characteristics by SCT status are shown in Table 1. Among 7743 participants, 583 had SCT and 4898 were women. Participants with SCT had a lower eGFR compared to those without SCT (86 vs. 90 ml/min/1.73 m2, p < 0.0001). The only other covariates to differ significantly by SCT were alcohol use and smoking status.

With median follow up of 7.1 years (range 0.4, 10.3 years), among the 583 participants with SCT, 85 (14-6%) experienced incident cognitive impairment, compared to 902 (11.7%) of those without SCT. Table 2 shows baseline characteristics by incident impairment on the SIS. Many baseline covariates had adverse levels in those with incident impairment, including age, eGFR, systolic blood pressure, hypertension, coronary heart disease, LVH, hyperlipidaemia and statin use. Those with incident impairment also had lower education and income at baseline, and were more likely to be men, than those without incident impairment.

In the univariate analysis, the OR of incident cognitive impairment by the SIS for those with versus without SCT was 1.18 (CI: 0.93, 1.51). Controlling for covariates, results were similar (OR 1.21; 95% CI: 0.94, 1.64). With addition of age, sex, education, income, region, eGFR, systolic blood pressure, diabetes, alcohol use, smoking status, exercise, coronary heart disease, left ventricular hypertrophy, atrial fibrillation, hyperlipidaemia, statin use, and hypertension. Note, y axis starts with a score of 5 words learning, memory, and semantic and phonemic fluency. These findings suggest that, unlike findings to date in SCA, biological consequences of SCT do not appear to cause cognitive dysfunction.

While there is literature suggestive that SCA patients experience cognitive dysfunction, to our knowledge, no prior studies evaluated SCT and cognitive impairment [5–9]. It is established that SCT is associated with hypercoagulability [25] and risk of both venous thromboembolism [19] and kidney disease [16,36,37]. In an autopsy series of 128 SCT patients, there appeared to be higher rates of visceral infections, including in the brain, than those without SCT (15% vs <1%) [22,38]. Additionally, SCT has been linked to mild cerebral vasculopathy in children on imaging [39]. Although specific mechanisms are unclear, and the findings are still debated, SCT may also be a risk factor for ischemic stroke, but not in the largest study of older adults to date, which included REGARDS participants [20–23]. Despite these findings, our results suggest that SCT does not lead to clinically significant cognitive dysfunction in adults aged 45 and older.

Regardless of the null findings here, and lack of differences in associations by age in this population age 45 and older, we would advocate for a detailed study examining cognitive impairment in a younger cohort of adults or in children. An estimated 11% of individuals with SCA experience a stroke before the age of 20, with the high risk period occurring between ages 2–5: approximately 24% experience stroke by age 45 [40,41]. Additionally, there are imaging findings suggestive of silent cerebroinfarction, including decreased brain volume [42] and increased white matter hyperintensities [8,9,11] in pediatric SCA patients. While there isn’t similar data on cognitive function, a meta-analysis on the impact of stroke and silent cerebral infarction on the intelligence quotient of patients with SCA indicated that those with prior stroke had an intelligence quotient 10 points lower than those with silent cerebral infarction, while those with silent cerebral infarction had an intelligence quotient 6 points lower than those without radiographic indications of

![Fig. 3. Sickle cell trait and longitudinal change in semantic fluency (Animal Fluency Test) and phonemic fluency (Letter F Test). Blue lines represent those with sickle cell trait and black lines those without sickle cell trait. Models were adjusted for age, sex, education, income, region, eGFR, systolic blood pressure, diabetes, alcohol use, smoking status, exercise, coronary heart disease, left ventricular hypertrophy, atrial fibrillation, hyperlipidaemia, statin use, and hypertension. Note, y axis starts with a score of 5 words learning, memory, and semantic and phonemic fluency. These findings suggest that, unlike findings to date in SCA, biological consequences of SCT do not appear to cause cognitive dysfunction. While there is literature suggestive that SCA patients experience cognitive dysfunction, to our knowledge, no prior studies evaluated SCT and cognitive impairment [5–9]. It is established that SCT is associated with hypercoagulability [25] and risk of both venous thromboembolism [19] and kidney disease [16,36,37]. In an autopsy series of 128 SCT patients, there appeared to be higher rates of visceral infections, including in the brain, than those without SCT (15% vs <1%) [22,38]. Additionally, SCT has been linked to mild cerebral vasculopathy in children on imaging [39]. Although specific mechanisms are unclear, and the findings are still debated, SCT may also be a risk factor for ischemic stroke, but not in the largest study of older adults to date, which included REGARDS participants [20–23]. Despite these findings, our results suggest that SCT does not lead to clinically significant cognitive dysfunction in adults aged 45 and older. Regardless of the null findings here, and lack of differences in associations by age in this population age 45 and older, we would advocate for a detailed study examining cognitive impairment in a younger cohort of adults or in children. An estimated 11% of individuals with SCA experience a stroke before the age of 20, with the high risk period occurring between ages 2–5: approximately 24% experience stroke by age 45 [40,41]. Additionally, there are imaging findings suggestive of silent cerebroinfarction, including decreased brain volume [42] and increased white matter hyperintensities [8,9,11] in pediatric SCA patients. While there isn’t similar data on cognitive function, a meta-analysis on the impact of stroke and silent cerebral infarction on the intelligence quotient of patients with SCA indicated that those with prior stroke had an intelligence quotient 10 points lower than those with silent cerebral infarction, while those with silent cerebral infarction had an intelligence quotient 6 points lower than those without radiographic indications of...](image)
silent cerebral infarction [43]. Given the evidence of early cognitive impairment and silent cerebral infarction in patients with SCA, it is possible that the childhood brain could be more susceptible to effects of SCT.

Limitations of this study should be considered. Cognitive function assessment ascertainment relied on participant contact by phone and incident cases may have been missed [26]. This issue was partly mitigated since we had a very large cohort, continuous decline in test scores could be examined with high precision and cohort retention was high with 87.3% cumulative retention from enrollment in 2003–7 to January 2011 [44]. Regardless, this type of bias would be predicted to bias results toward the null. Although SCT is less common among non-blacks, exclusion of other racial groups means that our findings may not be generalized to these groups. In order to minimize the impact of stroke itself on the associations of SCT with cognitive outcomes, we censored participants during follow up at the time of stroke. This could have contributed to the null findings we observed under an assumption that stroke is a mechanism whereby SCT might lead to cognitive impairment. However, this should not have been a factor in our findings since we recently reported no association of SCT with risk of stroke in REGARDS, a finding corroborated in three other cohorts including 19,000 participants [23]. It is possible that patients with SCT have undetected mild cognitive decline or subtle differences in parameters such as cognitive processing speed, that might be important to their functioning [10] and insensitive to the cognitive measures used. It is also possible that there is clinical relevance to the 20% increased odds of cognitive impairment based on the six-item screener, although this association was not statistically significant; the finding could be subject to type II error, but is consistent with the rest of the null findings. Other risk factors for cognitive impairment in REGARDS using this endpoint have had much higher relative risk estimates (e.g., 1.6 for male sex, 2.1 for black race and 2.2 for low education). Ongoing REGARDS research will classify participants on dementia status and re-analysis of sickle cell trait with this endpoint will be important. Finally, we did not include participants below age 45 or oversample very old people, groups where any impact of SCT on cognitive function might be different.

The strengths of this study include the prospective design and large geographically dispersed cohort of over 8000 black Americans with representation of men and women. Incident cognitive impairment and longitudinal cognitive performance were carefully determined through a variety of measures with robust null results across all of these. We included people aged 45 and older, the age group with the highest likelihood of cognitive impairment, allowing us a better possibility to detect an association if one exists. Indeed, over a median of 7.1 years of follow up we had nearly 1000 cases of incident cognitive impairment based on the SIS.

We present a detailed examination of the association of SCT with incident cognitive impairment and change in cognitive scores over time in a large population sample of blacks, showing no important association. The findings present are clinically relevant, as they might be reassuring to patients with SCT who are worried about cognitive function. The results suggest there is not cause for concern about cognitive dysfunction for SCT patients in this age group and their primary care providers. Further research is warranted to confirm our findings, particularly in younger individuals.

Declaration of Interest

The authors have no relevant relationships to disclose.

Author contributions

C.C., J.M.L., L.A.M., M.R.I., N.A.Z., R.N., F.U., V.G.W., S.E.J., J.M., H.H., C.W. and M.C. drafted/revised the manuscript for content. C.C., L.A.M., N.A.Z., R.N., F.U., V.G.W., S.E.J., C.W., H.H. and M.C. contributed to study design and concept. J.M.L. and L.A.M. performed statistical analysis.

C.C., J.M.L., L.A.M., M.R.I., N.A.Z., R.N., F.U., V.G.W., S.E.J., J.M., H.H., C.W. and M.C. interpreted data. C.W. contributed vital reagent for genotyping. F.U., V.G.W., S.E.J., C.W. and M.C acquired data. M.C. M.R.I., and R.N. supervised and coordinated the study. L.A.M., R.N., S.E.J., C.W. and M.C. obtained funding.

Acknowledgements

The authors thank other investigators, the staff, and the participants of the REGARDS study for their valuable contributions. A full list of participating REGARDS investigators and institutions can be found at http://www.regardsstudy.org. Study funding was by U01NS041588 from the National Institute of Neurological Disorders and Stroke, National Institutes of Health (NIH); American Society of Hematology HONORS Award; and HHSN26120080001E from the Intramural Research Program of the National Cancer Institute Center for Cancer Research, NIH. The content of this publication does not necessarily reflect the views or policies of the Department of Health and Human Services, nor does mention of trade names, commercial products, or organizations imply endorsement by the U.S. Government.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.eclinm.2019.05.003.

References

[1] Davíglus ML, Bell CC, Berteltini V, et al. National Institutes of Health State-of-the-Science Conference statement: preventing Alzheimer’s disease and cognitive decline. NIH Consens State Sci Statements 2010;27(4):1–30.
[2] Lorus N, Locascio JJ, Rentz DM, et al. Vascular disease and risk factors are associated with cognitive decline in the Alzheimer disease spectrum. Alz Dis Assoc Disord 2015;29(1):18–25.
[3] Duron E, Hanon O. Vascular risk factors, cognitive decline, and dementia. Vasc Health Risk Manag 2008;4(2):363–81.
[4] Blon K, Emmerin-Yonk MH, Koek HL. The influence of vascular risk factors on cognitive decline in patients with dementia: a systematic review. Maturitas 2013;76(2):113–7.
[5] Schatz J, Faiak RL, Kellett JM, Kramer JH. Cognitive functioning in children with sickle cell disease: a meta-analysis. J Pediatr Psychol 2002;27(8):739–48.
[6] Vichinsky EP, Neumayr LD, Gold JI, et al. Neuropsychological dysfunction and neuroimaging abnormalities in neurologically intact adults with sickle cell anemia. JAMA 2010;303(18):1823–31.
[7] Schatz J. Brief report: academic attainment in children with sickle cell disease. J Pediatr Psychol 2004;29(8):627–33.
[8] Pegelow CH, Macklin EA, Moser FG, et al. Longitudinal changes in brain magnetic resonance imaging findings in children with sickle cell disease. Blood 2002;99(8):1014–8.
[9] van der Land V, Hjimans CT, de Ruiter M, et al. Volume of white matter hyperintensities is an independent predictor of intellectual quotient and processing speed in children with sickle cell disease. Br J Haematol 2015;168(4):553–6.
[10] Jostesbury H, Kilkham FJ, Kolbe M, et al. White matter integrity and processing speed in sickle cell anemia. Neurology 2018;90(23):e2042–e50.
[11] Mackin RS, Iseal P, Turun D, et al. Neuroimaging abnormalities in adults with sickle cell anemia: associations with cognition. Neurology 2014;82(10):835–41.
[12] Torres LS, Okumura J, Silva DG, et al. Correction: inflammation in sickle cell disease: differential and down-expressed plasma levels of annexin A1 protein. PLoS One 2017;12(2):e0172859.
[13] Naik RP, Wilson JC, Eksunwe L, et al. Elevated D-dimer levels in African Americans with sickle cell trait. Blood 2016;127(18):2261–3.
[14] CoboC J, Allen JL. Hypoxemia in sickle cell disease: significance and management. Paediatr Respir Rev 2014;15(1):17–23.
[15] Sampietro M, Giovannetti T, Tarazi R. Hypoxia and inflammation in children with sickle cell disease: implications for hippocampal functioning and episodic memory. Neuropsych Rev 2014;24(2):252–65.
[16] Naik RP, Derebail VK, Grams ME, et al. Association of sickle cell trait with chronic kidney disease and albuminuria in African Americans. JAMA 2014;312(20):2115–25.
[17] Naik RP, Irvin MR, Judd S, et al. Sickle cell trait and the risk of ESRD in blacks. J Am Soc Nephrol 2017;28(7):2180–7.
[18] Austin H, Key NS, Benson JM, et al. Sickle cell trait and the risk of venous thromboembolism among blacks. Blood 2007;110(3):908–12.
[19] Folsom AR, Tang W, Roettler NS, et al. Prospective study of sickle cell trait and venous thromboembolism incidence. J Thromb Haemost 2015;13(1):2–9.
[20] Golomb MR. Sickle cell trait is a risk factor for early stroke. Arch Neurol 2005;62(11):1778–9.
[21] Dowling MM. Sickle cell trait is not a risk factor for stroke. Arch Neurol 2005;62(11):1780–1.
[22] Caughey MC, Loehr LR, Key NS, et al. Sickle cell trait and incident ischemic stroke in African Americans. JAMA Neurol 2018;75(7):802–7.
[23] Martin TW, Weisman IM, Zeballos RJ, Stephenson SR. Exercise and hypoxia increase sickling in venous blood from an exercising limb in individuals with sickle cell trait. Am J Med 1989;87(1):48–56.
[24] Westerman MP, Green D, Gilman-Sachs A, et al. Coagulation changes in individuals with sickle cell trait. Am J Hematol 2002;69(2):89–94.
[25] Howard VJ, Cushman M, Pulley L, et al. The Reasons for Geographic and Racial Differences in Stroke study: objectives and design. Neuroepidemiol 2005;25(3):135–43.
[26] Price AL, Patterson NJ, Plenge RM, Weinblatt ME, Shadick NA, Reich D. Principal components analysis corrects for stratification in genome-wide association studies. Nat Genet 2006;38(8):904–9.
[27] Levey AS, Stevens LA, Schmid CH, et al. A new equation to estimate glomerular filtration rate. Ann Intern Med 2009;150(9):604–12.
[28] Verduzco LA, Nathan DG. Sickle cell disease and stroke. Blood 2009;114(25):5117–25.
[29] Steen RG, Hankins GM, Xiong X, et al. Prospective brain imaging evaluation of children with sickle cell trait: initial observations. Radiology 2003;228(1):208–15.
[30] Ohene-Frempong K, Weiner SJ, Sleeper LA, et al. Cerebrovascular accidents in sickle cell disease: rates and risk factors. Blood 1998;91(1):288–94.
[31] Kawadler JM, Clayden JD, Clark CA, Kirkham FJ. Intelligence quotient in paediatric sickle cell disease: a systematic review and meta-analysis. Dev Med Child Neurol 2016;58(7):672–9.
[32] Cushman M, Callas PW, McClure LA, et al. N-terminal pro-B-type natriuretic peptide and risk of future cognitive impairment in the REGARDS cohort. J Alzheimers Dis 2016;54(2):497–503.