Cognitive Impairment Before Atrial Fibrillation–Related Ischemic Events: Neuroimaging and Prognostic Associations

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Background—It is likely that a proportion of poststroke cognitive impairment is sometimes attributable to unidentified prestroke decline; prestroke cognitive function is also clinically relevant because it is associated with poor functional outcomes, including death. We investigated the radiological and prognostic associations of preexisting cognitive impairment in patients with ischemic stroke or transient ischemic attack associated with atrial fibrillation.

Methods and Results—We included 1102 patients from the prospective multicenter observational CROMIS-2 (Clinical Relevance of Microbleeds in Stroke 2) atrial fibrillation study. Preexisting cognitive impairment was identified using the 16-item Informant Questionnaire for Cognitive Decline in the Elderly. Functional outcome was measured using the modified Rankin scale. Preexisting cognitive impairment was common (n=271; 24.6%). The presence of lacunes (odds ratio [OR], 1.50; 95% CI, 1.03–1.05; P=0.034), increasing periventricular white matter hyperintensity grade (per grade increase, OR, 1.38; 95% CI, 1.17–1.63; P<0.0001), deep white matter hyperintensity grade (per grade increase, OR, 1.26; 95% CI, 1.05–1.51; P=0.011), and medial temporal atrophy grade (per grade increase, OR, 1.63; 95% CI, 1.34–1.95; P<0.0001) were independently associated with preexisting cognitive impairment. Preexisting cognitive impairment was associated with poorer functional outcome at 24 months (mRS >2; adjusted OR, 2.43; 95% CI, 1.42–4.20; P=0.001).

Conclusions—Preexisting cognitive impairment in patients with atrial fibrillation–associated ischemic stroke or transient ischemic attack is common, and associated with imaging markers of cerebral small vessel disease and neurodegeneration, as well as with longer-term functional outcome.

Clinical Trial Registration—URL: http://www.clinicaltrials.gov. Unique identifier: NCT02513316. (J Am Heart Assoc. 2020;9:e014537. DOI: 10.1161/JAHA.119.014537.)

Key Words: atrial fibrillation • brain ischemia • cerebral small vessel disease • cognitive impairment • vascular dementia

P oststroke dementia is common, affecting up to 41.3% of patients in hospital populations.1 Atrial fibrillation (AF) is increasingly recognized as a key risk factor for dementia, both in association with and independent of clinically overt ischemic stroke.2 It is likely that a proportion of poststroke cognitive impairment is attributable to unidentified prestroke decline.1 The pooled prevalence of prestroke dementia is estimated to be 14.4% (in hospital-based cohorts, on the basis of data from 3 studies); it is associated with both neurodegenerative and vascular factors1,3 and with poor functional outcome.4–8 Most data on the clinical and radiological associations of prestroke cognitive impairment are from small, single-center studies in
Clinical Perspective

What Is New?

- We reviewed data from over 1000 patients presenting with ischemic stroke or transient ischemic attack associated with atrial fibrillation, recruited as part of the prospective, multicenter observational CROMIS-2 (Clinical Relevance of Microbleeds in Stroke 2) atrial fibrillation study.
- We found that preexisting cognitive impairment is common in patients presenting with atrial fibrillation–associated ischemic stroke or transient ischemic attack, affecting nearly a quarter of patients (24.6%).
- Preexisting cognitive impairment was associated with magnetic resonance imaging markers of cerebral small vessel disease (lacunes, white matter hyperintensities) and neurodegeneration (medial temporal atrophy), and was also associated with poorer functional outcome at 24 months, independent of the acute ischemic event.

What Are the Clinical Implications?

- Preexisting cognitive impairment might be underrecognized in patients presenting with ischemic stroke or transient ischemic attack associated with atrial fibrillation.
- It is important to identify preexisting cognitive impairment in these patients, as it appears to have prognostic implications.
- We used an informant questionnaire (the Informant Questionnaire for Cognitive Decline in the Elderly) to identify preexisting cognitive impairment, which might be a useful and relevant tool in acute stroke; further data validation of this tool is needed.

heterogeneous stroke populations, which might not be generalizable to AF-related stroke populations.\(^9\)–\(^19\) Most imaging studies have focused on global and regional atrophy measures and white matter changes, with limited descriptions of other important structural markers of small vessel disease (such as magnetic resonance imaging [MRI]-visible perivascular spaces and cerebral microbleeds).

We investigated the prevalence and associations of preexisting cognitive impairment in patients with ischemic stroke or transient ischemic attack (TIA) associated with AF. We hypothesized that patients with preexisting cognitive impairment would (1) have more evidence of small vessel disease and neurodegeneration that those without and (2) show associations with poorer functional outcome at 24 months.

Methods

Data Availability

Analyses for the CROMIS-2 (Clinical Relevance of Microbleeds in Stroke 2) study are ongoing; once all of these analyses are completed, the CROMIS-2 Steering Committee will consider applications from other researchers for access to anonymized source data.

Patient Selection

This is a predefined substudy nested within CROMIS-2 AF, a multicenter prospective observational study of patients with AF-related cardioembolic stroke or TIA, as described previously.\(^20,21\) Briefly, this was a study of adults (aged \( \geq 18 \) years) presenting with ischemic stroke or TIA with nonvalvular AF (confirmed by electrocardiography), who were eligible to start anticoagulation following their ischemic event.\(^20,21\) Patients who could not have an MRI scan, had contraindications to anticoagulation, or had previously received therapeutic anticoagulation, were excluded.\(^20,21\) The study was approved by the National Research Ethics Service (IRAS reference 10/H0716/61). Written informed consent was obtained in all cases.

Informant Questionnaire for Cognitive Decline in the Elderly

Preexisting cognitive impairment was identified using the 16-item Informant Questionnaire for Cognitive Decline in the Elderly (IQCODE). The informant (defined as the patient’s caregiver, family member, or friend) was asked to compare the patient’s cognitive and functional performance from 10 years before their stroke or TIA with their performance just before their stroke or TIA. The 16-item IQCODE includes 16 questions, each of which can be scored between 1 and 5; the total is then divided by 16 to provide the final score (range, 1.0–5.0). Preexisting cognitive impairment was defined as an IQCODE score \( > 3.3 \); this threshold was based on data from a systematic review evaluating the diagnostic accuracy of the IQCODE for detecting clinically diagnosed dementia (of any cause) in secondary care environments.\(^22\)

Preexisting cognitive impairment was defined as an IQCODE score \( > 3.3 \).\(^22\) All patients with a baseline IQCODE were included in this analysis.

Imaging

Imaging was performed acutely after the index ischemic event and completed locally at each study center in accordance with a standardized protocol, which has been published previously.\(^20\) Imaging was performed acutely after the index ischemic event and completed locally at each study center in accordance with a standardized protocol including axial T2, T2* gradient echo sequence, diffusion-weighted imaging, coronal T1, and fluid-attenuated inversion recovery (FLAIR) images.\(^20\) Sequence parameters were specified for T2*
gradient echo sequence; the remaining sequences were obtained according to local protocols.

All structural markers of cerebral small vessel disease were rated in accordance with consensus criteria, each measure was rated by a single individual blinded to all clinical information. Where possible, the hemisphere contralateral to the acute stroke was preferentially counted.

Previous cortical infarcts were identified using T2 and FLAIR sequences and confirmed as nonacute through comparison with diffusion-weighted images. Lacunes were identified and counted on T2 and FLAIR sequences using definitions from the Standards for Reporting and Imaging of Small Vessel Disease criteria. White matter hyperintensities (WMHs) in deep and periventricular distributions were rated on T2 and FLAIR sequences using the Fazekas scale. MRI-visible perivascular spaces in the basal ganglia and centrum semiovale were rated on T2 and FLAIR sequences using a previously described validated visual rating scale. Medial temporal atrophy (MTA) was rated on T1 or FLAIR coronal images using the Scheltens scale. Global cortical atrophy (GCA) was rated with the Pasquier scale using axial T1, FLAIR, or inverted T2 images. Cortical superficial siderosis was identified on T2* gradient echo sequences and classified as either focal, involving ≤3 sulci, or disseminated, involving ≥4 sulci. Cerebral microbleeds were rated using T2* gradient echo sequences using the Microbleed Anatomical Rating Scale.

Functional Outcomes Following the Index Ischemic Event

Functional outcome at 24 months was quantified using the modified Rankin scale (mRS) using multiple ascertainment methods to maximize follow-up; these included postal questionnaires sent to patients and their general practitioners, and death notifications from NHS Digital (previously the Health and Social Care Information Centre). The mRS was dichotomized, with a score of ≤2 indicating independence.

Statistical Analysis

We investigated for selection bias by comparing characteristics of those with and without a baseline IQCODE. We then compared baseline clinical, demographic, and imaging findings in patients with and without preexisting cognitive impairment. For all continuous variables, data were reviewed for normality, and if normally distributed, the independent t test was used. If variables were ordinal or not normally distributed, the nonparametric Mann–Whitney U test was used. Chi-squared or Fisher’s exact tests were used for categorical variables.

The results of univariable comparisons were used to identify variables for inclusion in multivariable logistic regression models; variables with P<0.20 were included in the adjusted analyses, except for situations where variables both described the same phenomenon (eg, clinical history of previous ischemic events and imaging evidence of a previous cortical infarct). The presence of ≥1 acute diffusion-weighted image lesions was included as a variable in all adjusted analyses for outcome, to control for the index event (ie, stroke or TIA). We adjusted for National Institutes of Health Stroke Scale (a clinical measure of stroke severity) and the presence of acute diffusion-weighted image changes as measures of the acute cerebral ischemic event; we did not additionally adjust for immediate postevent mRS for reasons of collinearity. We instead adjusted for preevent mRS to capture baseline function. Each model considered only a single neuroimaging marker at a time.

Post hoc analyses were performed after excluding those with a preexisting clinical diagnosis of dementia or cognitive impairment, previous ischemic events, or intracerebral hemorrhage at study entry. This was to establish whether any findings in the cohort as a whole were driven by patients with these diagnoses, which are associated with cognitive impairment.

Statistical analyses were performed (GB) using Stata (Version 15).

Results

Of the total number of patients included in the CROMIS-2 AF study (n=1490), we included 1102 patients for whom a baseline IQCODE was available; 388 patients were excluded as baseline IQCODE data were not available. The included patients were less likely to be current smokers (9.8% versus 16.9%; P=0.0001), more likely to have a formal diagnosis of dementia or cognitive impairment (2.8% versus 1.6%; P=0.166), less likely to have had a previous intracerebral hemorrhage (0.4% versus 1.1%; P=0.116), and more likely to be taking an antiplatelet drug at study entry (53.7% versus 48.7%; P=0.094). The included patients also had a slightly higher educational age (mean, 16.4 versus 16.9 years; P=0.027).

Baseline characteristics for our cohort are shown in Table 1. In our cohort (n=1102), the mean age was 76.0 years (SD, 10.1), and 471 patients (42.7%) were women.

Preexisting Cognitive Impairment and Associations

In our cohort, the mean IQCODE score was 3.2 (SD, 0.6; score range, 1.0–5.0), and 271 (24.6%) patients had IQCODE-
defined preexisting cognitive impairment, of whom 23 (8.5%) had a known diagnosis of dementia or cognitive impairment at study entry. When comparing baseline clinical and demographic characteristics (Table 1), those with IQCODE-defined preexisting cognitive impairment were older (79.2 years versus 75.4 years), more likely to be female (49.1% versus 40.7%), and have a diagnosis of hypertension (70.0% versus 60.6%), diabetes mellitus (20.7% versus 15.7%), heart failure (6.6% versus 3.6%), prior AF (37.7% versus 31.0%), and previous ischemic events (27.1% versus 16.5%). They were more likely to be taking an antiplatelet before their index event (62.1% versus 51.0%), and had a lower educational age (mean, 15.7 versus 16.6 years).

The neuroimaging features are presented in Table 2. Those with preexisting cognitive impairment were more likely to have previous cortical infarcts and lacunes. They had higher grades of periventricular WMHs, deep WMHs, MRI-visible perivascular spaces in the basal ganglia, MTA, and global cortical atrophy and were more likely to have multiple cerebral microbleeds. In multivariable logistic regression analysis (Table 3), in which each imaging predictor was considered separately, the presence of lacunes (OR, 1.50; 95% CI, 1.03–1.05), increasing periventricular WMHs (per grade increase, OR, 1.38; 95% CI, 1.17–1.63), deep WMHs (per grade increase, OR, 1.26; 95% CI, 1.05–1.51) and MTA (per grade increase, OR, 1.61; 95% CI, 1.34–1.95) grade were independently associated with preexisting cognitive impairment.

**Functional Outcome Data**

Outcome data at 24 months were available for 922 patients (83.7%) of whom 480 (52.1%) were functionally dependent (mRS >2). Preexisting cognitive impairment was associated with functional dependence at 24 months (n=157, 72.0% versus n=323, 45.9%) in univariable (unadjusted OR, 3.03; 95% CI, 2.18–4.23; P<0.0001), and multivariable analyses (OR, 2.43; 95% CI, 1.42–4.20; P<0.001), adjusted for age at event, sex, hypertension, hypercholesterolemia, diabetes mellitus, smoking, heart failure, clinical history of previous ischemic events, educational age, admission National Institutes of Health Stroke Scale, antiplatelet use, preevent mRS, and the presence of an acute diffusion-weighted image lesion at study entry.

**Subgroup Analyses**

We then repeated these analyses after excluding patients with a known clinical history of dementia, cognitive impairment, previous ischemic events, or intracerebral hemorrhage at study entry, to review whether the associations observed in the whole cohort were being driven by patients with these diagnoses (Tables S1 through S4). The results of univariable and multivariable associations with preexisting IQCODE were consistent with our main findings, except that the association with lacunes and preexisting cognitive impairment did not reach statistical significance in adjusted analyses.

**Discussion**

In our large, multicentre prospective cohort of patient with AF-associated ischemic stroke and TIA, we found that nearly a quarter of patients (24.6%) met IQCODE criteria for preexisting cognitive impairment; this was associated with the presence of lacunes, periventricular and deep WMHs, and medial temporal atrophy, but not with other structural markers of small vessel disease (MRI-visible perivascular spaces, cortical superficial siderosis, or cerebral microbleeds). We also found that IQCODE-defined cognitive impairment was associated with poorer functional outcome at 24 months.

Our findings in an AF-associated cohort are in keeping with previous studies that have shown that preexisting cognitive impairment is associated with both neurodegenerative and vascular factors. AF is increasingly recognized as a key risk factor for dementia, both in association with and independent of clinically overt ischemic stroke. Multiple mechanisms for this association have been proposed, including silent brain infarcts from recurrent embolization, cerebral hypoperfusion, chronic inflammation, and endothelial dysfunction or progression of preexisting cerebrovascular or neurodegenerative processes. We found rates of preexisting cognitive impairment that were higher than many unselected stroke populations, and our rates of impairment were also higher than those reported in other AF cohorts. This might reflect the variability in methods used to diagnose preexisting cognitive impairment, including different IQCODE thresholds.

Our finding that MTA is a common and prevalent finding in patients before stroke provides further evidence that this neuroimaging feature is important in AF-related ischemic stroke and TIA. AF has been shown to be associated with lower hippocampal volumes and poorer memory and learning performance in stroke-free individuals, and patients with AF have greater atrophy of their entorhinal cortex and medial temporal lobes, compared with those without. The relationship between AF and global atrophy measures is less clear; while one study found that AF was associated lower brain volumes globally, others did not identify an association, although this might reflect the younger age of these latter cohorts. An association with MTA but not global cortical atrophy is in keeping with our results and implicates Alzheimer disease pathology in the cognitive impairment associated with AF, a proposal for which there is supporting evidence.
longitudinal and pathological data.\textsuperscript{2,54–58} Proposed mechanisms by which AF might contribute to Alzheimer disease pathology include β- and γ-secretase inhibition, perivascular amyloid clearance failures and tau phosphorylation, all of which might be induced by AF-related cerebral hypoperfusion.\textsuperscript{2} However, MTA can also be a feature of vascular pathology, and it might be that this is the dominant pathology in AF.\textsuperscript{59,60}

We also found an association between WMH severity and preexisting cognition. Although WMHs in patients with AF might simply be attributable to age or a shared vascular risk factor profile,\textsuperscript{61} there is evidence to suggest that there is an independent association.\textsuperscript{62} While WMHs are associated with poorer cognitive performance,\textsuperscript{63} the data relating to cognitive impairment in AF and WMHs is conflicting, with some studies showing no association.\textsuperscript{50,51} We did not find an independent statistically significant association with imaging evidence of previous cortical infarcts, which might provide further evidence that embolism to the brain (either clinically overt or “silent”) is not the only mechanism contributing to cognitive impairment in these patients. The lack of association between preexisting cognitive impairment and other structural small vessel disease markers (MRI-visible perivascular spaces and cerebral microbleeds) is in keeping with data from other populations, which show inconsistent associations between cognitive impairment and these markers.\textsuperscript{64} The presence of both neurodegenerative and vascular pathologies support the argument that, in patients with ischemic stroke, preexisting dementia is a manifestation of “brain aging”\textsuperscript{3} rather than attributable to one single pathological process; this is in contrast with cognitive impairment before spontaneous intracerebral hemorrhage, where cerebral small vessel diseases (in particular, cerebral amyloid angiopathy) are important.\textsuperscript{19,65} This suggests that any future therapeutic strategies for cognitive impairment will need to be implemented early and before stroke to be effective.

We show that cognitive impairment before an ischemic event is important, as it influences subsequent functional outcome, independent of the acute ischemic event. This is in keeping with previous work from other centers, which has shown associations between prestroke dementia and poor functional outcomes.\textsuperscript{4–6} Appropriately identifying preexisting cognitive impairment can be important for decision making with regard to postevent rehabilitation.\textsuperscript{66} Questions remain about how best to diagnose preexisting cognitive impairment. The IQCODE has been used extensively\textsuperscript{67}; our data provide more evidence that the IQCODE might be a useful and relevant tool in acute stroke, as in our cohort it identifies patients at risk of subsequent cognitive impairment and poorer functional outcomes. While it might seem counterintuitive to not assess the patient directly, this is often useful in stroke where patients are unable to engage in formal testing, for example, because of aphasia or reduced consciousness. IQCODE-based estimates of cognition might prove more
Table 2. Comparison of Imaging Features Between Those With and Without Preexisting Cognitive Impairment

| Feature                                      | All          | Preexisting Cognitive Impairment | P Value |
|----------------------------------------------|--------------|----------------------------------|---------|
|                                              | Absent       | Present                          |         |
| n (%)                                        | 1102         | 831 (75.4)                       | 271 (24.6) | ... |
| Imaging evidence of previous cortical infarct | 207 (18.8)   | 142 (17.1)                       | 65 (24.1) | 0.011 |
| Lacunes, presence, n (%)                     | 188 (17.3)   | 130 (15.8)                       | 58 (22.1) | 0.020 |
| pvWMH grade, n (%)                           |              |                                  |         |
| 0                                            | 645 (58.5)   | 527 (63.4)                       | 118 (43.5) | <0.00001 |
| 1                                            | 206 (18.7)   | 149 (17.9)                       | 57 (21.0) |
| 2                                            | 195 (17.7)   | 125 (15.0)                       | 70 (25.8) |
| 3                                            | 56 (5.1)     | 30 (3.6)                         | 26 (9.6)  |
| dWMH grade, n (%)                            |              |                                  |         |
| 0                                            | 472 (42.8)   | 385 (46.3)                       | 87 (32.1) | <0.00001 |
| 1                                            | 431 (39.1)   | 315 (37.9)                       | 116 (42.8) |
| 2                                            | 129 (11.7)   | 94 (11.3)                        | 35 (12.9) |
| 3                                            | 70 (6.4)     | 37 (4.5)                         | 33 (12.2) |
| CSO-PVS grade, n (%)                         |              |                                  |         |
| 0                                            | 58 (5.4)     | 44 (5.4)                         | 14 (5.4)  | 0.5043 |
| 1                                            | 486 (45.2)   | 361 (44.3)                       | 125 (48.1) |
| 2                                            | 324 (30.1)   | 255 (31.3)                       | 69 (26.5) |
| 3                                            | 174 (16.2)   | 128 (15.7)                       | 46 (17.7) |
| 4                                            | 33 (3.1)     | 27 (3.3)                         | 6 (2.3)   |
| BG-PVS grade, n (%)                          |              |                                  |         |
| 0                                            | 70 (6.4)     | 54 (6.6)                         | 16 (6.0)  | 0.0033 |
| 1                                            | 782 (71.6)   | 607 (73.7)                       | 175 (65.3) |
| 2                                            | 183 (16.8)   | 130 (15.8)                       | 53 (19.8) |
| 3                                            | 52 (4.8)     | 30 (3.6)                         | 22 (8.2)  |
| 4                                            | 5 (0.5)      | 3 (0.4)                          | 2 (0.8)   |
| MTA grade, n (%)                             |              |                                  |         |
| 0                                            | 222 (22.0)   | 192 (24.9)                       | 30 (12.6) | <0.00001 |
| 1                                            | 470 (46.5)   | 373 (48.4)                       | 97 (40.6) |
| 2                                            | 229 (22.7)   | 161 (20.9)                       | 68 (28.5) |
| 3                                            | 66 (6.5)     | 38 (4.9)                         | 28 (11.7) |
| 4                                            | 23 (2.3)     | 7 (0.9)                          | 16 (6.7)  |
| GCA grade, n (%)                             |              |                                  |         |
| 0                                            | 355 (32.6)   | 282 (34.3)                       | 73 (27.3) | 0.0078 |
| 1                                            | 469 (43.1)   | 354 (43.1)                       | 115 (43.1) |
| 2                                            | 246 (22.6)   | 174 (21.2)                       | 72 (27.0) |
| 3                                            | 19 (1.7)     | 12 (1.5)                         | 7 (2.6)   |
| cSS, presence, n (%)                         | 3 (0.3)      | 1 (0.1)                          | 2 (0.7)   | 0.151 |
| CMB, presence, n (%)                         | 230 (20.9)   | 165 (19.9)                       | 65 (24.0) | 0.146 |
| Presence of >1 CMB, n (%)                    | 111 (10.1)   | 71 (8.5)                         | 40 (14.8) | 0.003 |

Percentage values were calculated using the total number of patients for whom data were available as the denominator. P values are from Mann–Whitney U tests (pvWMH, dWMH, CSO-PVS, BG-PVS, MTA, and GCA grades), Fisher’s exact test (cSS), or chi-squared tests (remainder). BG-PVS indicates magnetic resonance imaging–visible perivascular spaces in the basal ganglia; CMB, cerebral microbleed; CSO, magnetic resonance imaging–visible perivascular spaces in the centrum semiovale; cSS, cortical superficial siderosis; dWMH, deep white matter hyperintensities; GCA, global cortical atrophy; MTA, medial temporal atrophy; pvWMH, periventricular white matter hyperintensities.
Table 3. Multivariable Logistic Regression for Imaging Predictors of Preexisting Cognitive Impairment

| Imaging evidence of previous cortical infarct, presence* | OR  | 95% CI   | P Value |
|---------------------------------------------------------|-----|----------|---------|
| Lacunes, presence*                                       | 1.23| 0.84–1.78| 0.288   |
| pvWMH, per grade increase                                | 1.38| 1.03–1.05| 0.034   |
| dWMH, per grade increase                                 | 1.26| 1.05–1.51| 0.011   |
| BG-PVS, per grade increase                               | 1.16| 0.92–1.47| 0.212   |
| MTA, per grade increase                                  | 1.61| 1.34–1.95| <0.0001 |
| GCA, per grade increase                                  | 1.06| 0.86 to1.31| 0.588  |
| cSS, presence                                           | 8.21| 0.72–94.5| 0.091   |
| CMB, presence                                           | 1.10| 0.76–1.58| 0.620   |
| Presence of >1 CMB                                       | 1.49| 0.93–2.38| 0.093   |

Each model considered only a single neuroimaging marker at a time. BG-PVS indicates MRI-visible perivascular spaces in the basal ganglia; CMB, cerebral microbleed; cSS, cortical superficial siderosis; dWMH, deep white matter hyperintensities; GCA, global cortical atrophy; MTA, medial temporal atrophy; OR, odds ratio; pvWMH, periventricular white matter hyperintensities.

*Adjusted for age at event, sex, hypertension, diabetes mellitus, smoking, heart failure, known atrial fibrillation, educational age, and antiplatelet use. Remaining models were adjusted for age, sex, hypertension, diabetes mellitus, smoking, heart failure, clinical history of previous ischemic events, known atrial fibrillation, educational age, and antiplatelet use.

accurate than the potential overestimation resulting from acute patient testing (which can be influenced by intercurrent illness) and potential underestimation from formal dementia diagnoses. The association of IQCODE-defined cognitive impairment with recognized neurodegenerative and vascular neuroimaging markers suggests that this is indeed reflective of significant underlying pathology.

Strengths and Limitations

The strengths of this study include its large size and multicenter design. We considered a wide range of structural markers associated with cerebral small vessel disease and neurodegeneration. However, there are also some limitations. Patients without IQCODE data (and therefore excluded from our analysis) were more likely to have a formal diagnosis of dementia or cognitive impairment, suggesting some selection bias in our cohort. While the study imaging protocol required standardized sequences, there was still variability in how these sequences were obtained, as well as the MRI machines used by each center, and this could influence our imaging rating. This is particularly the case for cerebral microbleeds, where detection may have been influenced by magnetic resonance field strength; unfortunately, we did not have data on this. The mean age of our cohort is 76.0 years, and so these findings may not be generalizable to patients who have stroke at significantly younger ages; many of the brain pathologies that coexist and contribute to cognitive impairment are age related. We are unable to provide further data on whether patients had paroxysmal or persistent AF; there is some evidence that the former might have particular relevance for cognition in this context.\(^2,58,68\)

We do not have follow-up data on cardiac function for this cohort, and we acknowledge that this might have influenced functional outcome. Additionally, the CROMIS-2 AF study included only patients with recent cardioembolic ischemic stroke or TIA who were eligible for anticoagulation and able to undergo MRI scanning; this may not be representative of other patient groups. As we discussed above, MTA is not specific to Alzheimer disease; we do not have data on other markers of Alzheimer disease pathology (measures of Aβ and tau, e.g., using positron emission tomography or cerebrospinal fluid) and therefore cannot comment further on the extent to which pathologies associated with Alzheimer disease contribute to our findings. We also acknowledge that the IQCODE threshold used might not be equivalent to dementia and that a range of thresholds have been used in the past; the IQCODE has not been validated for prestroke impairment against a formal diagnosis of dementia, and we were unable to comment on this in our study. Formal neuropsychological testing of multiple domains would provide a more comprehensive assessment of cognition. Despite this, we would argue that cognitive impairment at the level identified by the IQCODE is still of relevance given that it is able to predict future outcomes in our cohort, although further validation is needed before the IQCODE can be used for this purpose in clinical practice.

Summary

In this comprehensive imaging description of the factors associated with preexisting cognitive impairment in
cardioembolic stroke and TIA, we report that preexisting cognitive impairment is common and associated with imaging markers of cerebral small vessel disease and neurodegeneration, as well as poorer functional outcomes at 24 months. We also provide evidence that the IQCODE might be useful as an acute tool in ischemic stroke and TIA, as it appears to identify those likely to have worse clinical outcomes. Future work validating the IQCODE in this context (including establishing the optimal threshold for these patients), together with further investigation of the factors that contribute to brain resilience and whether this can be influenced after ischemic injury, is needed.

Appendix
The CROMIS-2 Collaborators
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References
1. Pendlebury ST, Rothwell PM. Prevalence, incidence, and factors associated with pre-stroke and post-stroke dementia: a systematic review and meta-analysis. Lancet Neurol. 2009;8:1006–1018.
2. Ihara M, Washida K. Linking atrial fibrillation with Alzheimer’s disease: epidemiological, pathological, and mechanistic evidence. J Alzheimer’s Dis. 2018;62:61–72.
3. Pendlebury ST. Dementia in patients hospitalized with stroke: rates, time course, and clinico-pathologic factors. Int J Stroke. 2012;7:570–581.
4. Saposnik G, Cote R, Rochon PA, Mamdani M, Liu Y, Raptis S, Kapral MK, Black SE; Registry of the Canadian Stroke Network, Stroke Outcome Research Canada Working Group. Care and outcomes in patients with ischemic stroke with and without preexisting dementia. Neurology. 2011;77:1664–1673.
5. Wakisaka Y, Matsuo R, Hata J, Kuroda J, Kitazono T, Kamouchi M, Ago T; Fukuoka Stroke Registry Investigators. Adverse influence of pre-stroke dementia on short-term functional outcomes in patients with acute ischemic stroke: the Fukuoka Stroke Registry. Cerebrovasc Dis. 2017;43:82–89.
6. Melkas S, Oksala NK, Jokinen H, Pohjasvaara T, Vataja R, Oksala A, Kaste M, Karhunen PJ, Erkinjuntti T. Poststroke dementia predicts poor survival in long-term follow-up: influence of prestroke cognitive decline and previous stroke. J Neurol Neurosurg Psychiatry. 2009;80:845–870.
7. Henon H, Durieu I, Lebet F, Pasquier F, Leys D. Influence of prestroke dementia on early and delayed mortality in stroke patients. J Neurol. 2003;250:10–16.
8. Garcia-Ptacek S, Contreras Escamendez B, Zupanic E, Religa D, von Koch L, Johnell K, von Euler M, Kareholt I, Eriksdotter M. Prestroke mobility and dementia as predictors of stroke outcomes in patients over 65 years of age: a cohort study.
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Pasquier F, Henon H, Leys D. Relevance of white matter changes to pre- and poststroke dementia. Ann N Y Acad Sci. 2000;903:466–469.

Henon H, Pasquier F, Durieu I, Pruvo JP, Leys D. Medial temporal lobe atrophy and pre-existing dementia in stroke patients: relation to pre-existing dementia. Cognit Neuropsychol. 1998;5:641–647.

Pasquier F, Henon H, Leys D. Relevance of white matter changes to pre- and poststroke dementia. Ann N Y Acad Sci. 2000;903:466–469.

Pohjasvaara T, Mantyla R, Aronen HJ, Leskelä M, Salonen O, Kaste M, Erkinjuntti T. Clinical and radiological determinants of prestroke cognitive decline in a stroke cohort. J Neurol Neurosurg Psychiatry. 1999;67:742–748.

Barba R, Castro MD, del Mar Morin M, Rodriguez-Romero R, Rodriguez-García E, Canto R, Del Ser T. Prestroke dementia. Cerebrovasc Dis. 2001;11:216–224.

Caratossolo S, Riva M, Vicini Chilovi B, Cerea E, Mombelli G, Padovan A, Rozzini L. Prestroke dementia: characteristics and clinical features in a consecutive series of patients. Eur J Neurol. 2014;21:148–154.

Tan G, Chen SS, Chiu HF, Ungvari GS, Wong KS, Kwok TC, Mok V, Wong KT, Richards PS, Ahuja AT. Frequency and determinants of prestroke dementia in a Chinese cohort. J Neurol. 2004;251:604–608.

Horrstmann M, Rizos T, Rauch G, Fuchs M, Arden C, Veltkamp R. Atrial fibrillation and prestroke cognitive impairment in stroke. J Neurol. 2014;261:546–553.

Leitgeb C, Depienne D, Touze E, Henon H, Parretti L, Pasquier F, Gallau V, Leys D, Investigators SI. Prestroke dementia in patients with atrial fibrillation. Frequency and associated factors. J Neurol. 2005;252:1504–1509.

Banerjee G, Wilson D, Ambler G, Osei-Bonsu Appiah K, Shakeshaft C, Lunawat A, Mok VC, Wong A, Lam WW, Fan YH, Tang WK, Kwok T, Hui AC, Wong KS. Questionnaire on Cognitive Decline in the Elderly (IQCODE) for the diagnosis of dementia within a secondary care setting. Cochrane Database Syst Rev. 2015;10(suppl A100):155–161.

Chirdumio A, Wilson D, Ambler G, Osei-Bonsu Appiah K, Shakeshaft C, Brown MM, Al-Shahi Salman R, Jager HR, Werring DJ. CROMIS-2 Collaborators. Cognitive impairment before intracerebral hemorrhage is associated with cerebral amyloid angiopathy. Stroke. 2018;49:40–45.

Wilson D, Ambler G, Shakeshaft C, Brown MM, Chirdumio A, Al-Shahi Salman R, Lip G, Houlden H, Jager HR, Brown MM, Werring DJ. The Clinical Relevance of Microbleeds in Stroke study (CROMIS-2): rationale, design, and methods. Int J Stroke. 2015;10(suppl A100):155–161.

Chirdumio A, Wilson D, Shakeshaft C, Ambler G, White M, Cohen H, Youssy T, Lip GYH, Muir KW, Brown MM, Al-Shahi Salman R, Jager HR, Werring DJ. The Microbleed Anatomical Rating Scale (MARS): reliability of a tool to map brain microbleeds. Neurology. 2009;73:1759–1766.

Charidimou A, Linn J, Vernooij MW, Opherk C, Akoudad S, Baron JC, Greenberg SM, Jager HR, Werring DJ. The Microbleed Anatomical Rating Scale (MARS): reliability of a tool to map brain microbleeds. Neurology. 2009;73:1759–1766.

Optimizing cutoff scores for the Barthel index and the modified Rankin scale for defining outcome in acute stroke trials. Stroke. 2005;36:1984–1987.

Henon H, Pasquier F, Durieu I, Godefroy O, Lucas C, Lebert F, Leys D. Preexisting dementia in stroke patients. Baseline frequency, associated factors, and outcome. Stroke. 1997;28:2429–2436.

Kalantarian S, Stern TA, Mansour M, Ruskin JN. Cognitive impairment associated with atrial fibrillation: a meta-analysis. Ann Intern Med. 2013;158:338–346.

Chen LY, Norby FL, Gottesman RF, Mosley TH, Soliman EZ, Agarwal SK, Loehr LR, Folsom AR, Corsh J, Alonso A. Association of atrial fibrillation with cognitive decline and dementia over 20 years: the ARIC-NCS (Atherosclerosis Risk in Communities Neurocognitive Study). J Am Heart Assoc. 2018;7:e007301. DOI: 10.1161/JAHA.117.007301.

Kwok CS, Loke YK, Hale R, Potter JF, Mjint PK. Atrial fibrillation and incidence of dementia: a systematic review and meta-analysis. Neurology. 2011;76:914–922.

de Bruijn RF, Heerings J, Wolters FJ, Franco OH, Stricker BH, Hofman A, Koudstaal PJ, Ikram MA. Association between atrial fibrillation and dementia in the general population. JAMA Neurol. 2015;72:1288–1294.

Singh-Manoux A, Fayosse A, Sabia S, Canonico M, Bobak M, Elbaz A, Kivimaki M, Dugravot A. Atrial fibrillation as a risk factor for cognitive decline and dementia. Eur Heart J. 2017;38:2612–2618.

Santangelo P, Di Biase L, Bari M, Mohanty S, Pump A, Cereceda Brantes M, Horton R, Burkhardt JD, Lakkinenzy D, Reddy YM, Casella D, Lollo Russo A, Tondo C, Natale A. Atrial fibrillation and the risk of incident dementia: a meta-analysis. Heart Rhythm. 2012;9:1761–1768.

Gaita F, Corsinovi L, Anselmino M, Raimondo C, Panelli M, Toso E, Bergamasco L, Boffano C, Valentinis MC, Cesaroni F, Scaglione M. Prevalence of silent cerebral ischemia in paroxysmal and persistent atrial fibrillation and correlation with cognitive function. J Am Coll Cardiol. 2013;62:1990–1997.

Graff-Radford J, Madhavan M, Vemuri P, Rabinstein AA, Cha RH, Mielke MM, Kantarci K, Lowe V, Senjem ML, Gunter JL, Knopman DS, Petersen RC, Jack CR Jr, Roberts RO. Atrial fibrillation, cognitive impairment, and neuroimaging. Alzheimers Dement. 2016;12:391–398.

Chen LY, Lopez FL, Gottesman RF, Huxley RR, Agarwal SK, Loehr L, Mjint PK. Atrial fibrillation and cognitive decline-the role of subclinical cerebral infarcts: the Atherosclerosis Risk in Communities Study. Stroke. 2014;45:2568–2574.

Gardarsdottir M, Sigurdsson S, Aspelund T, Rokita H, Launer LJ, Gudnason V, Arnar DO. Atrial fibrillation is associated with decreased total cerebral blood flow and brain perfusion. Europace. 2018;20:1252–1258.

Lavy S, Stern S, Melamed E, Cooper G, Keren A, Levy P. Effect of chronic atrial fibrillation on regional cerebral blood flow. Stroke. 1980;11:35–38.

Tolato R, Corridoni C, Marinis C, Marsili R, Precipe M. Transcranial Doppler evaluation of cerebral blood flow in patients with paroxysmal atrial fibrillation. Ital J Neurol Sci. 1993;14:451–454.

Alosco ML, Spitznagel MB, Sweet LH, Josephson R, Hughes J, Gunstad J. Atrial fibrillation exacerbates cognitive dysfunction and cerebral perfusion in heart failure. Pacing Clin Electrophysiol. 2015;38:178–186.

Cacciator F, Testa G, Langellotta A, Galizia G, Della-Morte D, Gargiulo G, Bevilacqua A, Del Genio MT, Canonico V, Rengo F, Abete P. Role of ventricular rate response on dementia in cognitively impaired elderly subjects with atrial fibrillation: a 10-year study. J Neurol. 2012;213:143–148.

Hui DS, Morley JE, Mikolajczak PC, Lee R. Atrial fibrillation: a major risk factor for cognitive decline. Am Heart J. 2015;169:448–456.

Poggensi A, Inzitari D, Pantoni L. Atrial fibrillation and cognition: epidemiological data and possible mechanisms. Stroke. 2015;46:3316–3321.

Dietzel J, Haeusler KG, Endres M. Does atrial fibrillation cause cognitive decline and dementia? Europace. 2018;20:408–419.

Rivard L, Khairy P. Mechanisms, clinical significance, and prevention of cognitive impairment in patients with atrial fibrillation. J Cardiovasc Electrophysiol. 2017;28:1556–1564.

Knecht S, Oelschlager C, Duning T, Lohmann H, Albers J, Stehling C, Heindel W, Breithardt G, Berger K, Ringelstein EB, Kirchhof P, Wiersching H. Atrial fibrillation in stroke-free patients is associated with memory impairment and hippocampal atrophy. Eur Heart J. 2008;29:2125–2132.
51. Stefansdottir H, Amar DO, Aspelund T, Sigurdsson S, Jonsdottir MK, Hjaltason H, Launer LJ, Gudnason V. Atrial fibrillation is associated with reduced brain volume and cognitive function independent of cerebral infarcts. Stroke. 2013;44:1020–1025.

52. Qureshi AI, Saed A, Tasneem N, Adil MM. Neuroanatomical correlates of atrial fibrillation: a longitudinal MRI study. J Vasc Interv Neurol. 2014;7:18–23.

53. Seshadri S, Wolf PA, Beiser A, Elias MF, Au R, Kase CS, D'Agostino RB, DeCarli C. Stroke risk profile, brain volume, and cognitive function: the Framingham Offspring Study. Neurology. 2004;63:1591–1599.

54. Burton EJ, Barber R, Mukaetova-Ladinska EB, Robson J, Perry RH, Jaros E, Kalaria RN, O'Brien JT. Medial temporal lobe atrophy on MRI differentiates Alzheimer's disease from dementia with Lewy bodies and vascular cognitive impairment: a prospective study with pathological verification of diagnosis. Brain. 2009;132:195–203.

55. Rusanen M, Kivipelto M, Levalahti E, Laatikainen T, Tuomilehto J, Soininen H, Ngandu T. Heart diseases and long-term risk of dementia and Alzheimer's disease: a population-based CAIDE study. J Alzheimers Dis. 2014;42:183–191.

56. Dublin S, Anderson ML, Haneuse SJ, Heckbert SR, Crane PK, Breitner JC, McCormick W, Bowen JD, Teri L, McCurry SM, Larson EB. Atrial fibrillation and risk of dementia: a population-based cohort study. J Gerontol A Biol Sci Med Sci. 2014;69:609–615.

57. Bunch TJ, Weiss JP, Crandall BG, May HT, Bair TL, Osborn JS, Anderson JL, Muhlestein JB, Horne BD, Lappe DL, Day JD. Atrial fibrillation is independently associated with senile, vascular, and Alzheimer's disease. Heart Rhythm. 2010;7:433–437.

58. Dublin S, Anderson ML, Heckbert SR, Hubbard RA, Sonnen JA, Crane PK, Montine TJ, Larson EB. Neuropathologic changes associated with atrial fibrillation in a population-based autopsy cohort. J Gerontol A Biol Sci Med Sci. 2014;69:609–615.

59. Kalaria RN. The pathology and pathophysiology of vascular dementia. Neuropharmacology. 2018;134:226–239.

60. Bastos-Leite AJ, van der Flier WM, van Straaten EC, Steenekenbreg SS, Scheltens P, Barkhof F. The contribution of medial temporal lobe atrophy and vascular pathology to cognitive impairment in vascular dementia. Stroke. 2007;38:3182–3185.

61. Haesler KG, Wilson D, Fiebach JB, Kirchhoff P, Werring DJ. Brain MRI to personalise atrial fibrillation therapy: current evidence and perspectives. Heart. 2014;100:1408–1413.

62. Mayasi Y, Helenius J, McManus DD, Goddeau RP Jr, Jun-O’Connell AH, Moonis M, Henninger N. Atrial fibrillation is associated with anterior predominant white matter lesions in patients presenting with embolic stroke. J Neurol Neurosurg Psychiatry. 2018;89:6–13.

63. Prins ND, Scheltens P. White matter hyperintensities, cognitive impairment and dementia: an update. Nat Rev Neurol. 2015;11:157–165.

64. Banerjee G, Wilson D, Jager HR, Werring DJ. Novel imaging techniques in cerebral small vessel diseases and vascular cognitive impairment. Biochim Biophys Acta. 2016;1862:926–938.

65. Xiong L, Reijmer YD, Charidimou A, Cordonnier C, Viswanathan A. Intracerebrobral hemorrhage and cognitive impairment. Biochim Biophys Acta. 2016;1862:939–944.

66. Longley V, Peters S, Swarbrick C, Bowen A. What influences decisions about ongoing stroke rehabilitation for patients with pre-existing dementia or cognitive impairment: a qualitative study. Clin Rehabil. 2018;32:1133–1144.

67. McGovern A, Pendlebury ST, Mishra NK, Fan Y, Quinn TJ. Test accuracy of informant-based cognitive screening tests for diagnosis of dementia and multidomain cognitive impairment in stroke. Stroke. 2016;47:329–335.

68. Chen LY, Agarwal SK, Norby FL, Gottesman RF, Loeher LR, Soliman EZ, Mosley TH, Folsom AR, Coresh J, Alonso A. Persistent but not paroxysmal atrial fibrillation is independently associated with lower cognitive function: ARIC study. J Am Coll Cardiol. 2016;67:1379–1380.
SUPPLEMENTAL MATERIAL
|                                | All  | Pre-existing cognitive impairment | p value |
|--------------------------------|------|----------------------------------|---------|
|                                |      | Absent                           | Present |        |
| n (%)                          | 872  | 689 (79.0)                       | 183 (21.0) | <0.00001 |
| Age at event, years, mean (SD) | 75.1 (10.2) | 74.2 (10.2) | 78.5 (9.7) |         |
| Sex, female, n (%)             | 368 (42.2) | 280 (40.6) | 88 (40.1) | 0.070 |
| Hypertension, n (%)            | 519 (60.1) | 400 (58.6) | 119 (66.1) | 0.066 |
| Hypercholesterolaemia, n (%)   | 361 (41.9) | 286 (42.0) | 75 (41.7) | 0.936 |
| Diabetes mellitus, n (%)       | 137 (15.8) | 102 (14.9) | 35 (19.1) | 0.158 |
| Smoking at study entry, n (%)  | 91 (10.6) | 75 (11.0) | 16 (8.9) | 0.406 |
| Heart failure, n (%)           | 33 (3.8) | 21 (3.1) | 12 (6.6) | 0.027 |
| Known AF, n (%)                | 271 (31.4) | 206 (30.2) | 65 (35.9) | 0.138 |
| Educational age, years, mean (SD) | 16.5 (3.7) | 46.7 (3.2) | 15.7 (2.4) | 0.0031 |
| Admission NIHSS, median (IQR)  | 5 (2 to 10) | 5 (2 to 10) | 5 (2 to 10) | 0.9840 |
| Anti-platelet use, n (%)       | 395 (46.8) | 300 (44.9) | 95 (53.7) | 0.038 |

Comparison of baseline demographic and imaging characteristics between those with and without cognitive impairment prior to their qualifying event. Percentage values were calculated using the total number of patients for whom data was available as the denominator. p values are from independent t-tests (age, educational age), Mann-Whitney U test (NIHSS), Fisher’s exact test (previous intracerebral haemorrhage) or chi-squared tests (remainder). AF, atrial fibrillation; IQR, interquartile range; NIHSS, National Institutes of Health Stroke Scale; SD, standard deviation.
Table S2. Comparison of imaging features between those and without pre-existing cognitive impairment.

|                                | All     | Pre-existing cognitive impairment | p value |
|--------------------------------|---------|-----------------------------------|---------|
|                                | n (%)   | Absent   | Present  |         |
| n (%)                          | 872     | 689 (79.0) | 183 (21.0) | -       |
| Imaging evidence of previous cortical infarct, n (%) | 130 (14.9) | 93 (13.5) | 37 (20.3) | 0.021   |
| Lacunes, presence, n (%)      | 132 (15.4) | 97 (4.2)  | 35 (19.7)  | 0.073   |
| pvWMH grade, n (%)            |         |         |          | <0.0001 |
| 0                              | 531 (60.9) | 446 (64.7) | 85 (46.5) |         |
| 1                              | 166 (19.0) | 124 (18.0) | 42 (23.0) |         |
| 2                              | 141 (16.2) | 97 (14.1)  | 44 (24.0)  |         |
| 3                              | 34 (3.9)  | 22 (3.2)   | 12 (6.6)   |         |
| dWMH grade, n (%)             |         |         |          | <0.0001 |
| 0                              | 396 (45.4) | 337 (48.9) | 59 (32.2) |         |
| 1                              | 337 (38.7) | 253 (36.7) | 84 (45.9) |         |
| 2                              | 95 (10.9)  | 72 (10.5)  | 23 (12.6)  |         |
| 3                              | 44 (5.1)  | 27 (3.9)   | 17 (9.3)   |         |
| CSO-PVS grade, n (%)          |         |         |          | 0.9310  |
| 0                              | 50 (5.9)  | 38 (5.6)   | 12 (6.9)   |         |
| 1                              | 375 (44.0) | 298 (44.0) | 77 (44.0)  |         |
| 2                              | 261 (30.6) | 212 (31.3) | 49 (28.0)  |         |
| 3                              | 142 (16.7)| 111 (16.4) | 31 (17.7)  |         |
| 4                              | 24 (2.8)  | 18 (2.7)   | 6 (3.4)    |         |
| BG-PVS grade, n (%)           |         |         |          | 0.0422  |
| 0                              | 61 (7.1)  | 47 (6.9)   | 14 (7.8)   |         |
| 1                              | 624 (72.1)| 508 (74.2) | 116 (64.4) |         |
| 2                              | 141 (16.3)| 104 (15.2) | 37 (20.6)  |         |
| 3                              | 36 (4.2)  | 23 (3.4)   | 13 (7.2)   |         |
| 4                              | 3 (0.4)   | 3 (0.4)    | 0 (0.0)    |         |
| MTA grade, n (%)              |         |         |          | <0.0001 |
| 0                              | 193 (24.3)| 169 (26.6) | 24 (15.1)  |         |
| 1                              | 375 (47.2)| 311 (49.0) | 64 (40.3)  |         |
| 2                              | 162 (20.4)| 120 (18.9) | 42 (26.4)  |         |
| 3                              | 50 (6.3)  | 31 (4.9)   | 19 (12.0)  |         |
| 4                              | 14 (1.8)  | 4 (0.6)    | 10 (6.3)   |         |
| GCA grade, n (%)              |         |         |          | 0.106   |
| 0                              | 285 (33.1)| 236 (34.7) | 49 (27.2)  |         |
| 1                              | 378 (43.9)| 300 (44.1) | 78 (43.3)  |         |
| 2                              | 184 (21.4)| 137 (20.1) | 47 (26.1)  |         |
| 3                              | 14 (1.6)  | 8 (1.2)    | 6 (3.3)    |         |
| cSS, presence, n (%)          | 1 (0.1)  | 1 (0.2)    | 0 (0.0)    | 1.000   |
| CMB, presence, n (%)          | 173 (19.8)| 133 (19.3) | 40 (21.9)  | 0.441   |
| Presence of >1 CMB, n (%)     | 77 (8.8) | 55 (8.0)   | 22 (12.0)  | 0.087   |
Percentage values were calculated using the total number of patients for whom data was available as the denominator. p values are from Mann-Whitney U tests (pvWMH, dWMH, CSO-PVS, BG-PVS, MTA and GCA grades), Fisher’s exact test (cSS) or chi-squared tests (remainder). BG-PVS, MRI-visible perivascular spaces in the basal ganglia; CMB, cerebral microbleed; CSO, MRI-visible perivascular spaces in the centrum semi-ovale; cSS, cortical superficial siderosis; dWMH, deep white matter hyperintensities; GCA, global cortical atrophy; MTA, medial temporal atrophy; pvWVH, periventricular hyperintensities.
Table S3. Multivariable logistic regression for imaging predictors of pre-existing cognitive impairment.

| Imaging evidence of previous cortical infarct, presence | OR   | 95% CI        | p value |
|---------------------------------------------------------|------|---------------|---------|
| Lacunes, presence                                       | 1.47 | 0.94 to 2.31  | 0.093   |
| pvWMH, per grade increase                               | 1.32 | 1.08 to 1.61  | 0.006   |
| dWMH, per grade increase                                | 1.29 | 1.05 to 1.60  | 0.016   |
| BG-PVS, per grade increase                              | 1.03 | 0.77 to 1.36  | 0.854   |
| MTA, per grade increase                                 | 1.55 | 1.25 to 1.94  | <0.0001 |
| GCA, per grade increase                                 | 1.09 | 0.85 to 1.39  | 0.503   |
| CMB, presence                                           | 0.90 | 0.57 to 1.40  | 0.629   |
| Presence of >1 CMB                                      | 1.13 | 0.63 to 2.05  | 0.679   |

Each model considered only a single neuroimaging marker at a time. All remaining models were adjusted for age, sex, hypertension, diabetes mellitus, heart failure, known AF, educational age, and anti-platelet use. BG-PVS, MRI-visible perivascular spaces in the basal ganglia; CI, confidence interval; CMB, cerebral microbleed; dWMH, deep white matter hyperintensities; GCA, global cortical atrophy; MTA, medial temporal atrophy; OR, odds ratio; pvWMH, periventricular hyperintensities.
Table S4. Logistic regression models reviewing associations between IQCODE-defined pre-existing cognitive impairment and functional outcome at 24 months.

|                        | Univariable OR (95% CI) | p value  | Adjusted OR (95% CI) | p value  |
|------------------------|-------------------------|----------|----------------------|----------|
| Functional dependence  | 2.78 (1.88 to 4.10)     | <0.0001  | 3.33 (1.72 to 6.42)  | <0.0001  |
| (mRS > 2)              |                         |          |                      |          |

Multivariable model adjusted for age at event, sex, hypertension, hypercholesterolaemia, diabetes mellitus, smoking, heart failure, clinical history of previous ischaemic events, educational age, admission NIHSS, anti-platelet use, pre-event mRS and the presence of an acute DWI lesion at study entry. CI, confidence interval; DWI, diffusion weighted imaging; MoCA, Montreal Cognitive Assessment; mRS, modified Rankin scale; NIHSS, National Institutes of Health Stroke Scale.