Pattern and Rate of Cognitive Decline in Cerebral Small Vessel Disease: A Prospective Study

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Abstract

Objectives

Cognitive impairment, predominantly affecting processing speed and executive function, is an important consequence of cerebral small vessel disease (SVD). To date, few longitudinal studies of cognition in SVD have been conducted. We determined the pattern and rate of cognitive decline in SVD and used the results to determine sample size calculations for clinical trials of interventions reducing cognitive decline.

Methods

121 patients with MRI confirmed lacunar stroke and leukoaraiosis were enrolled into the prospective St George’s Cognition And Neuroimaging in Stroke (SCANS) study. Patients attended one baseline and three annual cognitive assessments providing 36 month follow-up data. Neuropsychological assessment comprised a battery of tests assessing working memory, long-term (episodic) memory, processing speed and executive function. We calculated annualized change in cognition for the 98 patients who completed at least two time-points.

Results

Task performance was heterogeneous, but significant cognitive decline was found for the executive function index (p<0.007). Working memory and processing speed decreased numerically, but not significantly. The executive function composite score would require the smallest samples sizes for a treatment trial with an aim of halting decline, but this would still require over 2,000 patients per arm to detect a 30% difference with power of 0.8 over a three year follow-up.
Conclusions
The pattern of cognitive decline seen in SVD over three years is consistent with the pattern of impairments at baseline. Rates of decline were slow and sample sizes would need to be large for clinical trials aimed at halting decline beyond initial diagnosis using cognitive scores as an outcome measure. This emphasizes the importance of more sensitive surrogate markers in this disease.

Introduction
Cerebral small vessel disease (SVD) is the most common cause of vascular cognitive impairment and vascular dementia[1,2]. Cross sectional studies have shown that the cognitive profile in SVD is characterized by early impairments of processing speed and executive function with a relative sparing of episodic memory[3–5]. These neuropsychological deficits have a significant impact and are associated with poor functional outcome, such as a reduction in instrumental activities of daily living[6]. A number of cross-sectional studies[3–5] have investigated the pattern of cognitive impairment in SVD, but longitudinal reports of how cognitive function changes over time is rare. This information is important for providing prognosis to patients with SVD, monitoring the progress of disease, and is particularly vital for planning treatment trials. Important issues include understanding the rate of cognitive change and identifying tasks which provide the most sensitive and reliable measures of cognitive change in this population.

SVD describes a heterogeneous condition ranging from mild asymptomatic white matter hyperintensities (WMH)[7] seen in community populations, through patients with isolated lacunar stroke, to individuals with multiple lacunar infarcts and WMH who may suffer from vascular dementia[1]. A younger onset monogenic form of SVD, Cerebral Autosomal-Dominant Arteriopathy with Ischemic Leukoencephalopathy (CADASIL), also occurs[1]. The rate of cognitive decline varies between these phenotypes and therefore careful description of the patient group is important in any study of SVD. Furthermore, many patients defined by radiological SVD, particularly older patients who present to memory clinics, may also have coexistent Alzheimer’s pathology. To study a group with relatively pure vascular cognitive impairment, we recruited patients presenting to a stroke service who, regardless of cognitive complaints, had radiologically confirmed lacunar infarcts and leukoaraiosis. Previous studies have shown that the cross-sectional presentation of cognitive impairment in this group is similar to that seen in CADASIL patients[3] in whom cognitive impairment occurs at a younger age when co-existent Alzheimer’s disease pathology is not present.

We determined the rate and pattern of cognitive change over a 3 year follow-up period in a prospective cohort of patients with clinical lacunar stroke and leukoaraiosis. We used this information to perform power calculations for intervention trials and determined which cognitive domains would be most sensitive to change.

Materials and Methods
Ethics Statement
Study protocols were approved by a local research ethics committee (London—Wandsworth) and all patients provided prior written informed consent. The study is registered with the UK clinical research network (http://public.ukcrn.org.uk/, study ID: 4577).
Participants

Participants were patients enrolled in the prospective St George’s Cognition And Neuroimaging In Stroke (SCANS) study[5,8,9], a longitudinal investigation into the relationship between MRI markers and cognition in patients with symptomatic SVD. For this study we used cognitive data acquired at baseline and annually during the first three years of follow-up. Patients were recruited from inpatient and outpatient stroke services of three hospitals in South London, UK (St George’s, King’s College and St Thomas’ Hospitals) between 2007 and 2010 and followed up annually with cognitive assessment and MRI. SVD was defined as a clinical lacunar stroke syndrome[10] with an anatomically appropriate lacunar infarct on MRI, in addition to confluent leukoaraiosis (Fazekas grade ≥2)[11]. Exclusion criteria were: 1) any stroke mechanism other than SVD (extra or intracranial large artery stenosis >50%, cardioembolic source, non-lacunar subcortical infarcts >1.5cm in diameter as these are often caused by emboli, or cortical infarcts); 2) a history of major neurological or psychiatric disorders (with the exception of depression)[5]; 3) non-fluent in English, and; 4) unwilling or unable to undergo MRI. Patients who suffered a subsequent clinical stroke remained in the study provided the new stroke was lacunar and met the inclusion criteria as above. All patients were studied at least three months after their most recent stroke to reduce influences of acute ischemia on cognition.

Neuropsychological Testing

Annually, patients underwent structured clinical examination, and completed a battery of widely used neuropsychological tasks chosen to characterize cognitive impairment in SVD [5,8]. Clinical assessment included the modified Rankin scale of disability and dependence following stroke[12]. Neuropsychological tasks are described in Table 1, and were grouped into four key cognitive domains. For each domain, task performance was evaluated on a common scale through the calculation of age-scale z-scores based on the best available published normative data[5,8] (see Table 1). Age scaling allows a meaningful average to be calculated within tasks related to a given domain. A Global Cognition measure of overall performance across all tasks was also produced by averaging all individual task scores. Parallel test forms were employed for two tests to reduce learning effects: the BMIPB Speed of Information Processing task (4 forms[13]) and single letter verbal fluency (annually alternating F-A-S and B-H-R). All other tasks were identical at each assessment. Premorbid IQ was measured using the National Adult Reading Test-Restandardized (NART-R)[14].

Cognitive change

Annualized change scores were computed to allow the analysis of patients with partial data and to account for variability in the timing of assessments. For each subject, and each task measure, a linear regression was fitted to the data: \( y = \alpha + \beta x \), where \( y \) is the scaled cognitive score and \( x \) the time (in years) from baseline. The estimated parameters \( \alpha \) and \( \beta \) therefore represent: the baseline score (\( \alpha \), the regression intercept) and the annualized change (\( \beta \), the regression slope). The latter can be viewed as an extension of the commonly employed difference score method (final score − initial score). Mean-average annualized change scores were computed across tasks with non-missing data in each cognitive domain (WM, LTM, PS, and EF). Exploratory data analysis confirmed that trends over this timescale were well described by a linear fit (see S1 Table).

Poor performance in neuropsychological testing. For two tasks (trail making and grooved pegboard; Table 1) raw performance is measured as the time taken to complete the task. To reduce skew and the impact of extreme scores, performances where the age-scaled
Z-score was less than $-3.33$ (corresponding to a scaled score $<0$) were recoded to this threshold value.

Floor effects, where a measure cannot discriminate performance below a certain point, are a particular concern for longitudinal studies of cognitive decline. When present, particularly at baseline, floor effects will reduce estimates of decline over time. To monitor this, we identified where subjects performed at the minimum possible level for each task, and by extension, cognitive indices where all the component tasks were performed at the minimum possible level.

### Sample size calculations

To assess implications for future research we produced 3-year difference scores (pro-rated from average annual change) and estimated the sample sizes required to detect treatment effects for both the cognitive indices and their constituent tasks. For example, if an average 3-year decline of 0.4 Z-units was observed on a measure, a treatment effect of 50% would be calculated based on the power to detect the difference between 0.4 in the control arm and 0.2 in the treatment arm given the observed standard deviation. We report sample size calculations based on 50, 40, 30 and 20 percent reductions in annual cognitive decline. Sample size calculations were based on a two-tailed, two-sample t-test with an alpha of 0.05 and a power of 0.80.

### Statistical Analysis

Analysis was carried out in R, version 3.02. Analyses were corrected for multiple comparisons using the Holm-Bonferroni method. Annualized cognitive change was tested using one-sample, two-tailed t-tests ($H_0$, cognitive change = 0). Effects of cognitive index on cognitive...
change were further investigated using a repeated measures ANOVA predicting cognitive change from the within-patients factor: cognitive index (levels: WM, LTM, PS and EF), with and without demographic covariates (age at baseline, gender, and NART-IQ). To investigate the influence of baseline task performance on cognitive change, tests for correlation (Pearson’s r) were conducted.

**Missing data.** Missing data where a complete session was lost are described in Fig 1. There were also some sporadic missing data for individual tasks (3.04% across 9 tasks). Such missing data occurred when parts of the task battery were not completed on a given occasion. Reasons included: time constraints, patient motivation, experimenter error, and other task-specific factors which made data unsuitable. Such missing data did not generally impact the cognitive change measure which uses average decline across tasks in a cognitive index. However, for two patients, subsequent to the baseline session, motor dexterity declined secondary to osteoarthritis (n = 1) and SVD-related disability (n = 1) such that they were unable to use a pen/pencil for fine motor control and, as a result, it was not possible to assess performance for any of the tasks comprising the processing speed index. These patients were excluded from all analyses of processing speed.

**Results**

**Demographics and patient flow**

There were 98 participants in whom cognition was obtained on at least two time-points and data from these participants are used in this analysis. This excluded participants without any follow-up (n = 22), and those without neuropsychological follow-up (n = 1). Most participants (n = 65) had all four time-points of data, 17 had 3 time-points and 16 had 2 time-points. Fig 1 presents a flow diagram of patient status at each time-point. The median inter-test interval was 1 year, 5.25 days, and the median deviation from the 12 month target was +5.75 days, (25th percentile: −2.25 days, 75th percentile: +25.75 days).

Three patients suffered a new clinical stroke during the study. Two were subcortical lacunar strokes and one a small cortical hemorrhage. The protocol specified that patients with recurrent lacunar stroke remain within the study, but one of the two patients was unable to continue due to disability. The cortical hemorrhage was considered a study endpoint. Two further patients met study endpoints by converting to vascular dementia, defined by DSM-IV criteria. Neither case was associated with new clinical stroke. One further patient suffered brain ischemia associated with a persisting global cognitive deficit following a cardiac arrest and was excluded from later time-points.

Stated reasons for withdrawal (n = 15) included unwillingness due to poor health (n = 7), unwilling/study not worthwhile (n = 5), could not tolerate psychology (n = 1), could not tolerate MRI (n = 1) and no reason given (n = 1). Deaths (n = 12), where cause could be ascertained (n = 10), were not due to SVD-stroke (cancer–n = 2, cardiovascular–n = 2, respiratory–n = 1, other (health related)–n = 3, other (health unrelated)–n = 2). Sample sizes at each assessment were reduced by sporadic missed sessions (n = 12) due to ill health, scheduling issues or non-attendance. Furthermore, some patients attended MRI but not cognitive testing (n = 3) and others completed only the MMSE from the cognitive test battery (n = 2).

Baseline demographic and risk factor information is shown in Table 2 for both those included in the follow-up and those in whom follow-up cognition data was not available. Patients lacking any follow-up data tended to be older, with more severe disability and significantly poorer baseline cognitive function (Table 2).
Fig 1. Flow diagram of study participant status. Middle column shows count data presented for total study membership; left shows valid neuropsychology assessment at each study time-point; and right are study exit events.

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Baseline Cognition

At baseline SVD patients displayed impairments predominantly in the PS and EF indices consistent with previous reports [5,8]. The WM and LTM indices did not differ significantly from normal performance (Table 3). Male gender (p < 0.02) and lower premorbid NART-IQ (p < 0.0001) were associated with lower performance, but increased age was not (p = 0.69).

Change in disability. There was a significant increase in disability, measured by the Rankin Scale [25], over the three year follow-up, with a mean (SD) annualised change of 0.43 (0.66; Table 3).

Cognitive Change

As a preliminary step, task data were plotted and the model fits inspected to confirm the suitability of the annualized change measure. We further estimated an objective measure of model
fit\textsuperscript{26,27} (the Akaike information criterion with small sample correction) to compare linear measures of change with quadratic. These results are presented in S1 Table.

Analysis of the profile of cognitive change showed that the annualized rate of change was not equal across cognitive domains ($F(4,368) = 6.13, p < 0.0001$; Fig 2). Executive Function exhibited a significant decline (Table 3 & Fig 2). WM, PS and Global cognition showed a non-significant reduction over time. The LTM index improved over time, most likely reflecting a practice or learning effect. Age, gender and premorbid IQ did not explain variability in cognitive change: age ($p = 0.99$); gender ($p = 0.66$); NART-IQ ($p = 0.83$). There were also no significant interactions between these variables and the cognitive index factor: index×age ($p = 0.82$); index×gender ($p = 0.41$); index×NART-IQ ($p = 0.81$). All correlations between baseline cognitive performance on a measure and subsequent rate of cognitive change for that measure were

### Table 3. Longitudinal analysis of cognition in SVD.

<table>
<thead>
<tr>
<th>Measure</th>
<th>Baseline</th>
<th>Annualized Change</th>
</tr>
</thead>
<tbody>
<tr>
<td>Working Memory</td>
<td>−0.13 (0.93), $p = 0.177$</td>
<td>−0.021 (0.37), $p = 0.57$</td>
</tr>
<tr>
<td>Episodic (Long Term) Memory</td>
<td>0.059 (0.97), $p = 0.55$</td>
<td>0.062 (0.26), $p = 0.02$</td>
</tr>
<tr>
<td>Processing Speed</td>
<td>−0.74 (0.89), $p &lt; 2 \times 10^{-12}$</td>
<td>−0.035 (0.23), $p = 0.13$</td>
</tr>
<tr>
<td>Executive Function</td>
<td>−0.77 (1.00), $p &lt; 8 \times 10^{-11}$</td>
<td>−0.09 (0.32), $p &lt; 0.007$</td>
</tr>
<tr>
<td>Global</td>
<td>−0.49 (0.82), $p &lt; 5 \times 10^{-8}$</td>
<td>−0.029 (0.20), $p = 0.15$</td>
</tr>
<tr>
<td>Rankin Scale Score</td>
<td>1.39 (0.93)</td>
<td>0.43 (0.66), $p &lt; 6 \times 10^{-9}$</td>
</tr>
</tbody>
</table>

Table presents Mean (SD), and $p$-value from one-sample t-tests for cognitive indices and clinical measures.

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**Fig 2. Estimates of annualized cognitive change cognitive indices.** Mean annualized cognitive change (filled circles) are displayed for the four key cognitive indices: Working Memory (WM), Episodic (Long-term) Memory (LTM), Processing Speed (PS) and Executive Function (EF). The composite Global change is also displayed. Error bars show the 95% confidence interval of the mean obtained from non-parametric bootstrap estimation. The dotted line represents stable performance. Significant change occurs where error bars do not cross the dotted line.

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non-significant: WM: $r = -0.23$, LTM: $r = 0.098$, PS: $r = -0.081$, EF: $r = -0.105$, Global: $r = 0.11$ (all $p > 0.1$), as were inter-index correlations (absolute $r$ ranged from 0.007 to 0.17; all $p > 0.1$). Individual-level cognitive change data are plotted (Fig 3) for the four cognitive indices. Performance was heterogeneous and some subjects showed stable performance over the study.

**Sample size calculations**

Table 4 compares the measures of cognition used in this study as potential endpoints in a future treatment study. Average 3-year difference scores are presented with the results of sample size calculations. The best performing individual tasks were single-letter verbal fluency (per arm $n = 2,776$, 30% effect) and the Wisconsin card sort task ($n = 3,524$, 30% effect). The multitask composite EF index performed better than any single task ($n = 2,192$, 30% effect vs. $n > 2,776$ for single tasks), an effect also seen in the PS composite ($n = 7,475$, 30% effect vs. $n > 9000$ for single tasks). This may be attributed to some of the favorable properties of multitask composite scores, which potentially allow performance to be assessed over a wider range of cognitive ability, and can diminish the impact of task-specific variability and ceiling/floor effects[28]. In this sample, floor level performances were commonly observed for several tasks, but especially the grooved pegboard (PS) and trail-making (EF) tests (GPT = 22.4%, TMT-B = 31.6%). In contrast, no subject performed at the minimum attainable level for any of the cognitive indices at baseline.

**Discussion**

During a three year follow-up study of cognition in patients with lacunar stroke and leukoaraiosis we report a pattern of change in cognition over time consistent with the baseline profile of impairments: average cognitive change declined most for the domains which were most impaired at baseline. However, cognitive change was variable and slow in most individuals such that measured declines were only statistically significant for executive function. The average rate of decline in executive function observed over 3-years in SVD is approximately a third of a standard deviation. This estimate in SVD is greater than comparable reports in healthy aging[29,30], but less than reported changes for general cognitive function, or memory scores, in both Alzheimer’s disease[31,32] and mild cognitive impairment[33].

Previous studies in SVD-related populations have investigated cognitive change in non-disabled individuals with radiological SVD (leukoaraiosis)[34], individuals with mixed vascular disease and vascular risk factors[35] and CADASIL[36,37]. In non-disabled individuals meeting radiological criteria for SVD[34] steeper 3-year cognitive decline than controls was reported for MMSE, verbal fluency (animal naming), Stroop color naming and trail making part A. No significant declines were found for immediate or delayed word recall, symbol digit modalities, digit span, trail making (B – A), or digit cancellation. Similarly, an older adult sample with vascular disease or risk factors[35] showed a 3-year decline on the symbol digit modalities test (comparable to the EF index in this sample), but no significant change in performance on a learning task. A large sample of CADASIL gene carriers followed up for an average of 24 months[36] showed significant decline in processing speed (trail making A) but not executive function (trail making B). In contrast, a small sample of CADASIL re-tested with a 7 year interval[37] showed significant decline on the CAMCOG battery (Global) and Stroop test (Executive), but not the trail-making task (Executive). In this group significant memory decline was also observed. Taken together with the results presented in our study from a clinically defined SVD population with radiological confirmation, there is significant task to task variability in decline over time in SVD, but more reliable decline in executive function tasks. It is an
Fig 3. Cognitive change for the four cognitive indices. Rows present data for the four cognitive indices, from the top: 1st row–Executive Function, 2nd row–Processing Speed, 3rd row–Working Memory, 4th row–Episodic (long-term) memory. Left figures present ‘spaghetti-plots’ where the cognitive index scores at each assessment (circles) are presented with scores for each subject joined by lines. Middle column figures present the lines of best fit from the linear model. Right column figures show the annualized change measure: the trajectory of change over time from the linear model.

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unanswered question whether the profile of decline is the same for CADASIL where samples span much larger age ranges.

Our results show that an executive function composite measure was the most sensitive to cognitive decline in this patient group, and that this was more sensitive than using individual cognitive tests. However, with any of the measures evaluated here, large sample sizes would be required to demonstrate the effectiveness of treatment aimed at halting cognitive decline. For example we estimate 2,192 patients in each arm would be required to detect a 30% reduction in the rate of EF cognitive decline during a three year follow-up. Even larger sample sizes would be required if other cognitive tests or domains were used. These numbers broadly agree with estimates from the LADIS study[38] where per-arm sample sizes required to detect a treatment effect over 3 years were estimated as 2,599 using the VADAS-cog battery and 1,809 using an EF composite measure. The large sample size estimates most likely reflect the slow and variable rate of decline in this patient group, although they may also be influenced by learning effects. Furthermore, looking at individual tasks, the better performing tasks were not those that SVD patients were most impaired on at baseline (trail-making, card sorting), but instead were tasks that showed smaller, but more reliable declines over the study period (single letter verbal fluency, digit symbol substitution). One explanation for this is the impact of performance floor effects for the former tests–individuals who perform at the lowest measurable level cannot subsequently decline regardless of their disease progress. These results suggest that future research into cognitive measures in SVD should focus on identifying executive function tasks which minimize practice effects and assess a wide range of deficit. However, the superior performance

<table>
<thead>
<tr>
<th>Index</th>
<th>Task</th>
<th>Baseline</th>
<th>3-years</th>
<th>Difference</th>
<th>50%</th>
<th>40%</th>
<th>30%</th>
<th>20%</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Mean (SD) Task Z-Scores</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Global</td>
<td>(All)</td>
<td>−0.5 (0.83)</td>
<td>−0.6 (1.1)</td>
<td>−0.097 (0.61)</td>
<td>2,492</td>
<td>3,893</td>
<td>6,921</td>
<td>15,571</td>
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<tr>
<td>WM</td>
<td>DS</td>
<td>−0.13 (0.93)</td>
<td>−0.19 (1.3)</td>
<td>−0.064 (1.1)</td>
<td>19,027</td>
<td>29,730</td>
<td>52,858</td>
<td>&gt;100k</td>
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<tr>
<td>LTM</td>
<td>Index</td>
<td>0.059 (0.97)</td>
<td>0.25 (1.3)</td>
<td>0.19 (0.78)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
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<tr>
<td></td>
<td>LogMem-I</td>
<td>0.19 (1.1)</td>
<td>0.34 (1.5)</td>
<td>0.15 (1.1)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
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<td></td>
<td>LogMem-D</td>
<td>0.35 (1.1)</td>
<td>0.69 (1.5)</td>
<td>0.35 (1.1)</td>
<td>-</td>
<td>-</td>
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<td>-</td>
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<tr>
<td></td>
<td>VisRep-I</td>
<td>−0.48 (1.2)</td>
<td>−0.3 (1.5)</td>
<td>0.15 (1.2)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
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<tr>
<td></td>
<td>VisRep-D</td>
<td>0.17 (1.1)</td>
<td>0.32 (1.5)</td>
<td>0.12 (1.2)</td>
<td>-</td>
<td>-</td>
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<td>-</td>
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<tr>
<td>PS</td>
<td>Index</td>
<td>−0.74 (0.89)</td>
<td>−0.8 (1)</td>
<td>−0.1 (0.69)</td>
<td>2,692</td>
<td>4,205</td>
<td>7,475</td>
<td>16,818</td>
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<td></td>
<td>SOIP</td>
<td>−0.85 (0.93)</td>
<td>−0.87 (1.1)</td>
<td>−0.054 (0.71)</td>
<td>10,803</td>
<td>16,879</td>
<td>30,006</td>
<td>67,512</td>
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<tr>
<td></td>
<td>GPT</td>
<td>−1.2 (1.4)</td>
<td>−1.3 (1.7)</td>
<td>−0.18 (1.3)</td>
<td>3,256</td>
<td>5,087</td>
<td>9,043</td>
<td>20,344</td>
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<tr>
<td></td>
<td>DSST</td>
<td>−0.13 (0.87)</td>
<td>−0.15 (0.97)</td>
<td>−0.067 (0.62)</td>
<td>5,434</td>
<td>8,489</td>
<td>15,092</td>
<td>33,955</td>
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<tr>
<td>EF</td>
<td>EF-Index</td>
<td>−0.77 (1.0)</td>
<td>−1.0 (1.3)</td>
<td>−0.27 (0.96)</td>
<td>790</td>
<td>1,234</td>
<td>2,192</td>
<td>4,931</td>
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<tr>
<td></td>
<td>SLVF</td>
<td>−0.2 (1.3)</td>
<td>−0.48 (1.6)</td>
<td>−0.28 (1.1)</td>
<td>1,000</td>
<td>1,562</td>
<td>2,776</td>
<td>6,245</td>
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<tr>
<td></td>
<td>TMT-B</td>
<td>−1.2 (1.6)</td>
<td>−1.4 (2.0)</td>
<td>−0.15 (1.6)</td>
<td>6,465</td>
<td>10,101</td>
<td>17,957</td>
<td>40,402</td>
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<tr>
<td></td>
<td>WCST</td>
<td>−0.9 (0.87)</td>
<td>−1.3 (1.6)</td>
<td>−0.38 (1.7)</td>
<td>1,269</td>
<td>1,983</td>
<td>3,524</td>
<td>7,928</td>
</tr>
</tbody>
</table>

Table presents mean (SD) Z-scores at baseline and 3-year follow-up and mean (SD) of their difference (follow-up – baseline) calculated per subject. Sample-size estimates show the number of patients per arm of a study required to detect a significant difference between groups equal to X% of the mean difference (alpha = 5%, power = 80%). Global—Global cognition index. WM—Working memory index. DS—Digit Span task. LogMem—Logical memory immediate (I) and delayed (D). VisRep—Visual reproduction task immediate (I) and delayed (D). LTM-Index—Long-term (Episodic) Memory Index. SOIP—BMIPB Speed of information processing task. GPT—Grooved pegboard task. DSST—Digit Symbol substitution task. PS-Index—Processing Speed index. SLVF—Single letter verbal fluency (FAS/BHR). TMT-B—Trail-making test-part B. WCST—Modified Wisconsin card sort test. EF-index—Executive function index.

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of composite measures coupled with the heterogeneity of change in SVD suggest that it may not be possible to have one single task to assess the condition. Finally, the large estimated sample sizes highlight the importance of alternative approaches to measuring cognitive function in treatment trials. One such approach is to use alternative markers to cognition, such as MRI measures[39] which have shown improved sensitivity to change in SVD[38]. A second approach is to improve sub-typing of SVD to identify those individuals likely to experience rapid cognitive decline.

This research should be interpreted with the following limitations in mind. First, no control group was included, so any cognitive decline could be caused by both SVD associated pathology and age related changes. However, previous case control studies have shown SVD is associated with impaired cognition compared with age matched normal control populations, and this has also been shown in the baseline data from the cohort we studied in this paper[5]. Second, sample size calculations were based on a hypothetical therapy which prevents cognitive decline subsequent to treatment. These calculations are not relevant where candidate therapies aim to treat existing cognitive impairments (i.e. to reverse cognitive decline), this is an aspect to be considered elsewhere.

In common with similar longitudinal studies[30,34,35,40], there was subject dropout, coupled with some systematic bias in the pattern of missing data. That is to say that despite considerable effort to assess all patients annually, a significant proportion did not complete the follow-up, and those who did not complete tended to be older, with greater disability and poorer cognition. Overall, the dropout rate was high (38%), and greater than similar studies of non-disabled elderly (e.g. LADIS[34]: 27%). As a result, the generalizability of our results may be reduced where substantially different patterns of withdrawal and/or loss to follow-up are obtained. This effect is likely to produce a modest underestimation of the average rate of cognitive change in SVD due to the effective censorship of data from those who decline the most, die, or withdraw from the research due to changed circumstances.

Furthermore, there are limitations related to the choice of cognitive measures. We selected measures that are widely used and that have previously been employed to investigate cognition in SVD[34,40,41], and we grouped task performance into cognitive domains. However neuropsychological tasks often lack complete specificity to any one cognitive domain and results should be consequently be interpreted with caution. Particularly, not all tasks controlled for motor impairment. As a result, some tasks contributing to the executive function, processing speed and episodic memory indices have significant motor performance elements rather than pure cognitive effects. Affected tasks include trail making, grooved pegboard, visual reproduction and digit symbol substitution. However, the decline seen in executive function factor is unlikely to be better explained by worsening motor speed deficits in SVD as the non-motor verbal fluency and card sorting tasks show larger declines. There were also practice effects present in the psychological data, most noticeably in the tasks assessing episodic memory where average performance increased over time. It is unclear the degree to which practice effects were present in other tasks and cognitive domains, however, where present, practice effects would also act to underestimate the degree of cognitive decline. The use of tasks with alternate forms, particularly for memory-based tasks would reduce this effect and so this is both a limitation of this research and a recommendation for future research. The learning effect for memory tasks highlights the preservation of memory in SVD and contrasts with the prominent decline reported in the AD/MCI phenotype[32,42]. Finally, cognitive trajectory is a simplistic model of change over time, but one that suits the characteristics of this data where task measures and inter-subject slopes are variable, and the participants who drop out tend to show the largest changes. With longer follow-up, more complex models of cognitive decline in SVD may be informative and particularly may be able to capture non-linear aspects of change.
In conclusion, in a group of patients with symptomatic SVD we found a significant decline in performance on executive function tasks over three years. In contrast, working memory and processing speed did not decline significantly, and practice effects were seen for episodic memory. An analysis of the sample size required for a treatment study established that large sample sizes would be needed to demonstrate the effectiveness of a treatment which acts to slow the rate of cognitive decline in SVD. These findings emphasise the importance of surrogate disease markers in SVD such as those provided by MRI.

Supporting Information

S1 Table. Descriptive statistics for Akaike information criterion. Descriptive statistics for the Akaike information criterion with small sample correction (AICc) are presented for linear and quadratic change in cognition for participants with complete data over all time points (n = 64). Smaller AICc values indicate superior fit. Mean average AICc values favor a linear fit over quadratic given the data.

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Author Contributions

Conceived and designed the experiments: AJL TRB RGM HSM. Performed the experiments: AJL EAZ TRB. Analyzed the data: AJL RLB EAZ TRB RGM HSM. Contributed reagents/materials/analysis tools: AJL RGM HSM. Wrote the paper: AJL RLB EAZ TRB RGM HSM.

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