Small vessel disease lesion type and brain atrophy: The role of co-occurring amyloid

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Abstract

Introduction: It is unknown whether different types of small vessel disease (SVD), differentially relate to brain atrophy and if co-occurring Alzheimer's disease pathology affects this relation.

Methods: In 725 memory clinic patients with SVD (mean age 67 ± 8 years, 48% female) we compared brain volumes of those with moderate/severe white matter hyperintensities (WMHs; n = 326), lacunes (n = 132) and cerebral microbleeds (n = 321) to a reference group with mild WMHs (n = 197), also considering cerebrospinal fluid (CSF) amyloid status in a subset of patients (n = 488).

Results: WMHs and lacunes, but not cerebral microbleeds, were associated with smaller gray matter (GM) volumes. In analyses stratified by CSF amyloid status, WMHs and lacunes were associated with smaller total brain and GM volumes only in amyloid-negative patients. SVD-related atrophy was most evident in frontal (cortical) GM, again predominantly in amyloid-negative patients.

Discussion: Amyloid status modifies the differential relation between SVD lesion type and brain atrophy in memory clinic patients.

KEYWORDS
Alzheimer’s disease, brain atrophy, cerebral microbleeds, cerebral small vessel disease, lacunes, magnetic resonance imaging, vascular cognitive impairment, white matter hyperintensities

1 INTRODUCTION

Brain atrophy is common in memory clinic patients and is related to cognitive decline.1,2 Yet understanding of the etiology of brain atrophy in these patients is incomplete. Evidently, Alzheimer’s disease (AD) and other neurodegenerative pathologies are key contributors to atrophy, but a role of vascular brain injury, in particular cerebral small vessel disease (SVD), is also increasingly recognized.3-5 SVD is an etiological heterogeneous construct. The most common forms are arteriolosclerosis and cerebral amyloid angiopathy.6 Different forms of SVD manifest themselves in different types of vascular brain lesions, which can be seen on magnetic resonance imaging (MRI):
white matter hyperintensities of presumed vascular origin (WMHs), lacunes, and cerebral microbleeds.\(^2\) It is unclear whether different MRI manifestations of SVD differentially relate to brain atrophy. Previous studies have mostly focused on a single lesion type, in particular WMHs, without directly comparing different lesion types. Yet different lesion types can reflect different disease processes in different parts of the brain, potentially with different relationships with gray matter (GM) degeneration.\(^7\) Evaluation of such potential differences may provide further leads on disease processes underlying atrophy.

A substantial proportion of memory clinic patients with SVD will also have AD,\(^8,9\) but insight into the interplay between vascular brain injury and AD pathological processes is incomplete. Large studies assessing both processes in relation to brain atrophy are needed, as was recently stressed.\(^4,10\) We therefore investigated the relation between different SVD lesion types on MRI and brain atrophy (in terms of severity and pattern) in a large population of memory clinic patients and determined whether this relation was modified by cerebrospinal fluid (CSF) amyloid biomarker status.

2 | MATERIALS AND METHODS

2.1 | Study population

Patients were included from the TRACE-VCI study cohort, which consists of 860 memory clinic patients from two Dutch tertiary outpatient clinics (Amsterdam University Medical Center and University Medical Center Utrecht). For the TRACE-VCI study, patients were eligible if their MRI showed at least a minimal burden of vascular brain injury, including:

- mild WMHs (Fazekas scale grade 1\(^11\)) and presence of \(\geq 2\) vascular risk factors (hypertension, hypercholesterolemia, diabetes mellitus, obesity, current smoking, or a reported history of a vascular event other than stroke);
- moderate/severe WMHs (Fazekas \(\geq 2\));
- \(\geq 1\) lacune(s);
- \(\geq 1\) (sub)cortical infarct(s);
- \(\geq 1\) cerebral microbleed(s);
- \(\geq 1\) intracerebral hemorrhage(s).\(^12\)

Patients were included regardless of severity of their cognitive deficit, including patients with no objective cognitive impairment, mild cognitive impairment (MCI), and dementia. Patients with a primary non-vascular and non-neurodegenerative etiology (eg, brain tumor, hydrocephalus, or excessive alcohol consumption) were excluded. For the current study on SVD, we excluded patients with non-lacunar (sub)cortical infarcts (\(n = 96\)) and intracerebral hemorrhages (\(n = 11\)), because presence of these lesions can substantially affect brain volumes. Of the remaining 753 patients, brain volume measurements could be generated in 725 patients, which comprise the current study population (see Appendix B, Figure SB1 in supporting information).

The study was approved by the institutional review boards of the Amsterdam University Medical Center and the University Medical Center Utrecht. All subjects provided written informed consent prior to any research-related procedures.

2.2 | Image acquisition

Patients were scanned on one of six different MRI scanners: four at the Amsterdam University Medical Center and two at the University Medical Center Utrecht. MRI vendors were either General Electric (GE) or Philips. All patients were scanned using an MRI protocol that included a 3D T1-weighted, fluid-attenuated inversion recovery (FLAIR) and T2*-weighted/susceptibility-weighted imaging sequence ([12]; for details see Appendix A.1 in supporting information).

2.3 | Cerebral SVD lesion types on MRI

Presence of WMHs (using the Fazekas scale for deep WMHs, ranging from 0 to 3\(^11\)), lacunes, and cerebral microbleeds was rated on FLAIR and T2*-susceptibility weighted images by a resident neurologist (RH, 4 years of experience) and a neuroradiologist (in training; JdB, 11 years of experience), using the STRIVE (standards for reporting vascular changes in neuroimaging) criteria.\(^3\)

2.4 | WMH and brain volume measurements

We specifically used segmentation tools known to be relatively insensitive to interscanner differences and a high burden of pathology.\(^13\) A semi-automated processing pipeline was used to obtain WMH and brain volumes (for details see Appendix A.2 in supporting information). This pipeline consisted of (1) automatic WMH segmentation using k-nearest neighbor classification with tissue type priors; (2) lesion filling using the SLF-toolbox; (3) automatic segmentation of lesion-filled images using the Computational Anatomic Toolbox (CAT12). Lacunes, non-lacunar infarcts, intracerebral hemorrhages, and incidental findings were segmented manually using an in-house developed MeVisLab (MeVis Medical Solutions AG, Bremen, Germany) tool.\(^14-16\)

Total brain volume (TBV) was defined as the sum of GM and white matter (WM) volumes (including the volume of pathology). Regional GM, WM, and CSF volumes were obtained using the Hammers atlas in CAT12, dividing the brain into 68 regions of interest: 32 per hemisphere and four infratentorial regions [http://brain-development.org].\(^17-20\) To compensate for variability in head size, all brain volumes were normalized using the “residual normalization method.”\(^21\)

2.5 | Cerebrospinal fluid testing

CSF concentrations of amyloid beta\(_{42}\) (or A\(_{42}\)), tau, and/or total tau phosphorylated at threonine 181 (or p-tau) were measured using...
commercially available enzyme-linked immunosorbent assays (ELISAs) at a central laboratory for clinics at the Department of Clinical Chemistry of the Amsterdam University Medical Center22 (for details see Appendix A.3 in supporting information). Patients with a low CSF concentration of $A_\beta_{42} (< 640 \text{ ng/L})$ were assumed to have a high brain amyloid load, and hence were considered amyloid positive.23 For the present study, CSF data were available for 488 patients (67%).

2.6 Statistical analysis

All analyses were performed using IBM SPSS version 25 unless specified (for details see Appendix A.4 in supporting information).

2.6.1 Relation between lesion type and brain atrophy

The relation between lesion type and brain atrophy was explored with linear regression analyses (to obtain an indication of effect sizes) as well as with Bayesian network analyses (to investigate independent relations between different lesion types and brain atrophy). In the linear regression analysis, TBV, GM, and WM volumes (normalized for head size) were compared between each lesion type and a reference group with only Fazekas score 1 and ≥ 2 vascular risk factors (but no other lesions). To limit the effect of using different MR scanners on brain volume measurements, we included a model term for scanner type (using effect coding). We also adjusted for age, sex, and additionally for clinical diagnosis (no objective cognitive impairment, MCI, or dementia) and the number of vascular risk factors. We also used log-transformed WMH volume as a continuous variable.

Bayesian network analyses (bnlearn R package24; for settings see Duering et al.25) were used to assess the conditional dependencies between each lesion type, age, and sex (potential determinants) and brain volumes (outcome). In Bayesian network analyses, variables with a deterministic influence on the outcome (conditionally independent variables) are identified and separated from others with only an indirect influence on the outcome. Networks are produced in which direct determinants are connected directly to the outcome, while conditionally independent variables are connected only indirectly, via other variables. Normalized brain volumes were used as outcome (standardized into $z$-scores using the mean and standard deviation [SD] of the whole study sample and corrected for scanner effect). The strength of the connections between direct determinants and outcome was assessed by 100 bootstrap replications. Only connections that were strong enough (> 40% replications) were shown in the final network.

For WMH, which proved to be the strongest determinant of atrophy, we also assessed if it was associated with atrophy in specific brain regions. Normalized regional brain volumes were standardized into $z$-scores using the mean and SD of the whole study sample and corrected for age, sex, and scanner effect. We compared the relationship between log WMH volume and regional brain volumes for all 68 regions using linear regression analyses (with Bonferroni correction for multiple testing, adjusting for age, sex, and scanner effect).

2.6.2 Modification by co-occurring amyloid pathology

To investigate whether the relation between lesion type and brain atrophy was modified by co-occurring amyloid pathology, we assessed possible interactions between each lesion type and CSF amyloid status in relation to brain atrophy. We also performed post-hoc analyses stratified for CSF amyloid status. Normalized brain volumes were standardized in the same manner as described before, but using the mean and SD of the CSF subgroup, amyloid-negative or amyloid-positive subgroup depending on the specific analysis.

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**TABLE 1** Baseline characteristics of the total study population

<table>
<thead>
<tr>
<th>Number of patients, n total = 725</th>
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</thead>
<tbody>
<tr>
<td><strong>Demographics</strong></td>
</tr>
<tr>
<td>Age, years</td>
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<tr>
<td>Sex, female</td>
</tr>
<tr>
<td>Years of education</td>
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<tr>
<td><strong>Vascular risk factors</strong></td>
</tr>
<tr>
<td>Hypertension</td>
</tr>
<tr>
<td>Hypercholesterolemia</td>
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<tr>
<td>Diabetes mellitus</td>
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<tr>
<td>Current smoker</td>
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<tr>
<td>Obesity, body mass index ≥ 30</td>
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<tr>
<td>History of stroke</td>
</tr>
<tr>
<td>History of reported vascular events other than stroke</td>
</tr>
<tr>
<td><strong>Number of vascular risk factors</strong></td>
</tr>
<tr>
<td><strong>Clinical diagnosis</strong></td>
</tr>
<tr>
<td>No objective cognitive impairment</td>
</tr>
<tr>
<td>Mild cognitive impairment</td>
</tr>
<tr>
<td>Dementia</td>
</tr>
<tr>
<td>Vascular</td>
</tr>
<tr>
<td>Neurodegenerative</td>
</tr>
<tr>
<td>Alzheimer’s disease</td>
</tr>
<tr>
<td>Frontotemporal disease</td>
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<tr>
<td>Lewy body dementia</td>
</tr>
<tr>
<td>Others</td>
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<tr>
<td>Unknown etiology</td>
</tr>
</tbody>
</table>

Notes: Data are presented as mean ± standard deviation, number of patients (percentage of total study population) or median [interquartile range]. For years of education data were present in 720 patients. For vascular risk factors, current smoker and obesity data were present in 718 and 713 patients, respectively. aSuch as primary progressive aphasia, cortical basal syndrome, and progressive supranuclear palsy. bDementia of unknown origin, further examination needed to state diagnosis.
FIGURE 1  Occurrence of lesion types. Venn diagram showing the occurrence of lesion types in the entire study population (n = 725) as well as in the cerebrospinal fluid (CSF) amyloid-positive (n = 261) and amyloid-negative (n = 227) patients in the CSF subgroup. In 719 patients (99%), information regarding presence/absence of cerebral microbleeds (CMBs) was present. The number of patients with a certain lesion type (alone or in combination with another lesion type) is shown. The colors represent the percentage of the respective patient group, illustrating which (combination of) lesion types were observed. The majority of patients only had mild white matter hyperintensities (WMHs; Fazekas score of 1) or moderate/severe WMHs (Fazekas score 2 or 3) but no other lesions. Multiple lesion types occurred in 321 patients (44%) of the entire study population. Of 382 patients (53%) with either cerebral microbleeds/lacunes, 242 (63%) had multiple cerebral microbleeds/lacunes; 71 patients (10%) had multiple lacunes (max: 30). 171 patients (24%) had multiple cerebral microbleeds (CMBs; max: ~500). Of the patients with CMBs, 37 patients (12%) had only deep CMBs, 212 patients (66%) had only lobar CMBs, and 70 patients (22%) both had deep and lobar CMBs. In two patients, no information regarding CMB location was available.

2.7 | Data availability

The data that support the findings of this study are available from the corresponding author, upon reasonable request.

3 | RESULTS

3.1 | Baseline characteristics

The 725 patients had a mean (± SD) age of 67 (± 8) years and 348 (48%) were female. Baseline characteristics are shown in Table 1. A total of one thousand seventy six patients (24%) had no objective cognitive impairment, 175 (24%) MCI, and 374 (52%) dementia. Baseline characteristics of patients with available CSF (n = 488, 67%) are shown in Appendix C, Table CS1 in supporting information. Patients with available CSF were younger (t = 2.8, P = .006), less often had hypertension (χ² = 8.6, P = .003) or hypercholesterolemia (χ² = 5.7, P = .017), and were less often obese (χ² = 4.2, P = .04) compared to those without available CSF. In the CSF subgroup, amyloid-negative patients were younger (t = -4.9, P < .001) and more often obese (χ² = 8.9, P = .003) than amyloid-positive patients. Amyloid-negative patients more often had no objective cognitive impairment (χ² = 72.3, P < .001) and were less often demented (χ² = 52.9, P < .001) than amyloid-positive patients.

The occurrence of each lesion type (and the overlap between lesion types) is shown in Figure 1. A total of three thousand twenty one patients (44%) had multiple lesion types. Moderate/severe WMHs (Fazekas score 2 or 3) were present in 326 patients (45%), lacunes in 132 patients (18%), and cerebral microbleeds in 321 patients (44%).
TABLE 2  Relationship between lesion type and brain volumes for total study population and stratified for CSF amyloid biomarker status

<table>
<thead>
<tr>
<th></th>
<th>Total brain volume</th>
<th>Gray matter volume</th>
<th>White matter volume</th>
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<tbody>
<tr>
<td>Total study population (n = 725)</td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Mild WMHs and ≥ 2 VRF (reference; n = 197)</td>
<td>—</td>
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<td>—</td>
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<tr>
<td>Moderate/severe WMHs (n = 326)</td>
<td>–0.02 [–0.10;0.05]</td>
<td>–0.07 [–0.14;–0.002]</td>
<td>0.05 [–0.03;0.13]</td>
</tr>
<tr>
<td>Lacunes (n = 132)</td>
<td>–0.06 [–0.15;0.03]</td>
<td>–0.08 [–0.17;0.004]</td>
<td>0.0001 [–0.10;0.10]</td>
</tr>
<tr>
<td>Cerebral microbleeds (n = 321)</td>
<td>–0.02 [–0.09;0.06]</td>
<td>–0.05 [–0.12;0.03]</td>
<td>0.03 [–0.05;0.11]</td>
</tr>
<tr>
<td>CSF amyloid negative (n = 227)</td>
<td></td>
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<tr>
<td>Mild WMHs and ≥ 2 VRF (reference, n = 71)</td>
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<tr>
<td>Moderate/severe WMHs (n = 88)</td>
<td>–0.10 [–0.24;0.04]</td>
<td>–0.14 [–0.27;–0.01]</td>
<td>–0.002 [–0.15;0.14]</td>
</tr>
<tr>
<td>Lacunes (n = 42)</td>
<td>–0.22 [–0.37;–0.07]</td>
<td>–0.22 [–0.37;–0.08]</td>
<td>–0.10 [–0.27;0.07]</td>
</tr>
<tr>
<td>Cerebral microbleeds (n = 88)</td>
<td>–0.09 [–0.23;0.05]</td>
<td>–0.07 [–0.21;0.06]</td>
<td>–0.06 [–0.21;0.08]</td>
</tr>
<tr>
<td>CSF amyloid positive (n = 261)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mild WMHs and ≥ 2 VRF (reference; n = 62)</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Moderate/severe WMHs (n = 126)</td>
<td>0.08 [–0.05;0.22]</td>
<td>0.05 [–0.08;0.18]</td>
<td>0.08 [–0.06;0.22]</td>
</tr>
<tr>
<td>Lacunes (n = 42)</td>
<td>0.08 [–0.10;0.26]</td>
<td>0.07 [–0.10;0.25]</td>
<td>0.04 [–0.14;0.23]</td>
</tr>
<tr>
<td>Cerebral microbleeds (n = 130)</td>
<td>0.12 [–0.02;0.25]</td>
<td>0.10 [–0.03;0.23]</td>
<td>0.08 [–0.06;0.21]</td>
</tr>
</tbody>
</table>

Notes: Data are presented as standardized beta coefficients with 95% confidence intervals after correction for age, sex, and scanner effect. All brain volumes were corrected for variations in head size using the total intracranial volume. Abbreviations: CSF, cerebrospinal fluid; VRF, vascular risk factors; WMH, white matter hyperintensities.

of which 37 (12%) had only deep, 212 (66%), only lobar and 70 patients (22%) both deep and lobar cerebral microbleeds (CMBs). (In two patients, information on CMB presence/location was missing). The reference group of only Fazekas score 1 and ≥ 2 vascular risk factors (but no other lesions) consisted of 197 patients (27%). In the CSF subgroup, amyloid-positive patients more often had moderate/severe WMHs compared to amyloid-negative patients (χ² = 4.5, P = .04). Occurrence of lacunes and cerebral microbleeds did not differ between amyloid-negative and amyloid-positive patients (see Figure 1).

In the entire study population, mean TBV (normalized for head size, in mL) was 1042 ± 60 mL, mean GM volume 565 ± 44 mL, and mean WM volume 477 ± 33 mL. Median WMH volume was 6 mL (range 0.05 to 166 mL). In the CSF subgroup, amyloid-positive patients had a smaller TBV (amylloid positive: 1035 ± 24 mL; amyloid negative: 1064 ± 30 mL; standardized β coefficient [95% CI], adjusted for age, sex, and scanner effect: –0.19 [–0.27;–0.11], P < .001) and GM volume (amyloid positive: 558 ± 20 mL; amyloid negative: 585 ± 23 mL; β: –0.26 [–0.33;–0.18], P < .001), but similar WM volume (amyloid positive: 476 ± 14 mL; amyloid negative: 479 ± 15 mL; β: –0.01 [–0.09;0.08], P = .86) compared to the amyloid-negative patients.

3.2 | WMHs relate to GM atrophy

WMHs, in particular, were related to brain atrophy: compared to the reference group (Fazekas score 1), patients with moderate/severe WMHs (Fazekas score 2–3) had smaller GM volume (standardized β coefficient [95% confidence interval (CI)]: –0.07 [–0.14;–0.002], P = .045; additionally adjusted for clinical diagnosis: –0.06 [–0.12;0.005], P = .07), but TBV and WM volumes did not differ significantly (Table 2). When WMH volume was entered as a continuous variable, it was also significantly associated with TBV (β: –0.03) as well as GM volume (β: –0.06;0.07), but not WM volume. Patients with cerebral microbleeds had no significant differences in brain volumes compared to the reference group, although a trend with smaller GM volume was observed (β: –0.08 [–0.15;0.004], P = .06; additionally adjusted for clinical diagnosis: –0.008 [–0.16;0.007], P = .03). Patients with cerebral microbleeds had no significant differences in brain volumes compared to the reference group. This was also the case for patients with multiple (≥ 5), lobar (pure or any), or deep microbleeds. Adjusting for the number of vascular risk factors did not change any of the previous results. There was no significant interaction between clinical diagnosis and moderate/severe WMHs, lacunes, or microbleeds in relation to TBV, GM, or WM volume (interaction terms all P > .05).

The Bayesian network analyses integrating all lesion types confirmed the relation between WMHs and GM volume (see Figure 2). These analyses showed WMHs directly determined GM volume, independent of lacunes and cerebral microbleeds.

3.2.1 | Pattern of SVD-related brain atrophy

Brain atrophy in relation to WMHs occurred in specific brain regions (see Figure 3). Differences were most evident in frontotemporal cortical regions, but also the postcentral gyrus, thalamus, and anterior
3.3 Relation between WMHs and GM atrophy is modified by amyloid status

Analyses in patients with available CSF (n = 488) showed that the relation between lesion type and brain atrophy was influenced by co-occurring amyloid pathology (Table 2). There was a significant inverse interaction between CSF amyloid status and lesion type in relation to brain atrophy (interaction term WMH burden × CSF amyloid status for TBV: $P = .012$; GM volume: $P = .006$), lacunes (TBV: $P = .012$; GM volume: $P = .006$), and cerebral microbleeds (TBV: $P = .01$; GM volume: $P = .02$). In amyloid-negative patients, moderate/severe WMHs were associated with smaller GM volume ($-0.14 [-0.27; -0.01]$, $P = .035$) and lacunes with smaller TBV ($-0.22 [-0.37; -0.07]$, $P = .004$) and GM volume ($-0.22 [-0.37; -0.08]$, $P = .003$) compared to the reference group. By contrast, in amyloid-positive patients, none of the small vessel lesion types was associated with brain atrophy (Table 2). Vice versa, the effect size of the reduction in TBV and GM volume associated with a positive compared to negative CSF amyloid status was almost twice as large in patients with a low WMH burden relative to patients with a high WMH burden (Appendix C, Table CS3 in supporting information). The same was observed for the relation between the CSF amyloid status and TBV and GM volume in the absence versus the presence of lacunes.

The Bayesian network analyses confirmed the relation between lesion type and brain atrophy in amyloid-negative patients (data not shown), with lacunes directly determining GM volume. Additionally, WMHs had an indirect effect (via lacunes) on GM volume (data not shown).

The pattern of SVD-related brain atrophy also differed depending on CSF amyloid status (Figure 4). In amyloid-negative patients, right mid frontal gyrus, left and right thalamus, and right precentral gyrus, GM volumes were significantly smaller in patients with high WMH burden compared to patients with low WMH burden (Bonferroni-corrected $P < .05$, see Appendix C, Table CS4 in supporting information). In amyloid-positive patients, only left superior temporal gyrus GM volumes were significantly smaller in patients with high WMH burden compared to patients with low WMH burden (Bonferroni-corrected $P < .05$, Appendix C, Table CS4 in supporting information). No significant interaction between WMH burden and CSF amyloid status was found after correcting for multiple testing (Bonferroni-corrected $P > .05$). Regional brain volume analyses were not performed for patients with and without lacunes, due to a lack of power.

4 DISCUSSION

This study demonstrates that different manifestations of SVD differentially relate to brain atrophy in memory clinic patients, especially in the absence of concurrent amyloid pathology. In particular WMHs and lacunes were related to brain atrophy, WMH associated brain atrophy was most pronounced in frontal cortical GM regions.
Regional brain volume analysis. Effect size map showing the relation between log white matter hyperintensity (WMH) volume and regional gray matter (GM) volume using standardized beta coefficients (red: GM volume smaller in patients with higher log WMH volume; blue: GM volume smaller in patients with lower log WMH volume) in all patients (n = 725). Across all patients higher log WMH volume was associated with smaller GM volume in several (predominantly frontotemporal) brain regions. * Bonferroni-corrected P < .05

4.2 Pattern of SVD-related brain atrophy

Previous studies that investigated the pattern of SVD-related brain atrophy found a relation between WMHs and predominantly cortical GM atrophy.\textsuperscript{34–36,38–40} Regarding lacunes, most studies also found a relation with cortical GM atrophy,\textsuperscript{35,38–40} while one study also found an association with subcortical GM atrophy.\textsuperscript{40}

Only three of the previous studies\textsuperscript{34,35,39} have specifically investigated the regional pattern of cortical GM atrophy. Two studies\textsuperscript{34,39} were done in patients with symptomatic lacunes with moderate/severe WMHs, while the other\textsuperscript{35} used memory clinic patients with a history of stroke due to SVD or a lacune on MRI. All three studies found a rather global pattern of cortical GM atrophy in relation to WMHs, with relative sparing of hippocampal and medial temporal regions. In the present study, investigating a large cohort of memory clinic patients with SVD, GM atrophy in relation to WMHs was predominantly seen in frontotemporal, but not parietal-occipital, regions.

4.3 Modification by amyloid pathology

The interplay between SVD and AD pathological processes has gained increasing attention in the past decade.\textsuperscript{41} While many studies have investigated the interaction between amyloid and WMHs in relation to cognition and prognosis, with variable results (see Roseborough et al.\textsuperscript{42}), brain atrophy as an outcome has hardly been studied.\textsuperscript{43,44} We now show that there is indeed an interaction between WMHs and CSF amyloid status in relation to brain atrophy. WMHs and lacunes were related to atrophy, but primarily in amyloid-negative patients. Vice versa, the relation between CSF amyloid status and brain atrophy was much stronger in patients with a low burden of WMH or lacunes.

There is thus converging evidence that both SVD and AD pathological processes contribute to brain atrophy in memory clinic patients, but for each this is most evident in the absence of the other. This could perhaps be explained by a ceiling effect, possibly linked to disease stage. Longitudinal studies could give more insight into the combined effects of WMHs and amyloid beta on brain atrophy over time and reveal the temporality of both disease processes.
FIGURE 4 Regional brain volume analysis in cerebrospinal fluid (CSF) subgroup. Effect size map showing the relation between log white matter hyperintensity (WMH) volume and regional gray matter (GM) volume using standardized beta coefficients (Red: GM volume smaller in patients with higher log WMH volume; blue: GM volume smaller in patients with lower log WMH volume). A, CSF amyloid-negative patients (n = 261); B, CSF amyloid-positive patients (n = 227). The stratified analyses show that the effect is highly dependent on CSF amyloid status. While amyloid-positive patients have a lower GM volume than amyloid-negative patients, the association between higher log WMH volume and more GM atrophy was more pronounced in several brain regions for amyloid-negative patients only. * Bonferroni-corrected $P < .05$
4.4 | “Distant” effects of SVD

The exact mechanism by which SVD contributes to brain atrophy is not yet clear. It has been suggested SVD can lead to secondary degeneration: a process in which cortical GM atrophy occurs as a result of disconnection of WM tracts.33,36,40,45–47 This “disconnection phenomenon” has been demonstrated in patients with cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy and ischemic stroke.47,48 The underlying cellular mechanisms behind this process are not clear yet.49 Our results showed primarily subcortical lesions (WMHs and lacunes) were related with predominantly frontal cortical GM atrophy. This “distant” cortical atrophy in relation to these subcortical vascular lesions may indeed reflect secondary neurodegeneration through disconnection. The predominant involvement of frontal brain regions may be explained by increased susceptibility of axons in frontal regions to impaired cerebral perfusion and vascular injury, leading to partial frontal lobe disconnection.50

4.5 | Strengths and limitations

Strengths of our study include the large sample size, and detailed information on brain volumes and all SVD lesion types. Furthermore, CSF was available in a substantial subset of patients, allowing us to assess the impact of SVD both in the absence and presence of biomarker evidence of concomitant amyloid pathology.

Some limitations need to be addressed. First, selection bias could play a role because patients were included at a tertiary memory clinic and not all patients underwent a lumbar puncture. Second, all patients in the present study had some degree of vascular brain injury (as this was part of the inclusion criteria for the TRACE-VCI study), which could have led to an underestimation of the observed effects. Nevertheless, there was a great variability in the burden of vascular brain injury, which should have allowed us to detect relevant effects. Fourth, we used multicenter MRI data from different scanners. However, we used a high-quality, semi-automatic segmentation pipeline (including lesion filling) and corrected for scanner effect in our analyses. Both amyloid-positive and -negative patients were scanned using the same variety of MRI scanners instead of a single (different) scanner for each subgroup. Therefore the differences found in these patients cannot be explained by a scanner effect. Finally, the cross-sectional design of the present study does not allow us to infer causality regarding SVD and brain atrophy.

5 | CONCLUSION

In this study, we demonstrate that different manifestations of SVD related differentially to brain atrophy in memory clinic patients. In particular WMHs and lacunes were related to brain atrophy, but mainly in the absence of concurrent amyloid pathology. Vice versa, the relation of AD pathological processes and brain atrophy is most evident when the burden of SVD is low. These findings show that the most common types of brain pathologies in people attending a memory clinic, SVD and AD pathological processes contribute to atrophy, likely through different pathways.

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CONFLICTS OF INTEREST

There are no conflicts of interest for any of the authors.

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**SUPPORTING INFORMATION**
Additional supporting information may be found online in the Supporting Information section at the end of the article.

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