Frequency and Location of Dilated Virchow-Robin Spaces in Elderly People: A Population-Based 3D MR Imaging Study

**BACKGROUND AND PURPOSE:** dVRS have been previously associated with aging and cerebrovascular diseases. However, little is known about their prevalence and topographic distribution in the general elderly population.

**MATERIALS AND METHODS:** dVRS were evaluated by using high-resolution 3D MR imaging in 1826 subjects enrolled in the 3C-Dijon MR imaging study. On T1-weighted MR imaging, dVRS were detected according to 3D imaging criteria and rated by using 4-level severity scores based in the BG or in the WM. The number and anatomic location of large dVRS (>3 mm) were recorded.

**RESULTS:** dVRS were observed in the BG or WM in every subject. The severity of dVRS was significantly associated with higher age in both the BG and WM, whereas sex was related to the severity of dVRS only in the BG. Large dVRS were detected in 33.2% of participants. Status cribrorum was found in 1.3% of participants. dVRS were also highly prevalent within the hippocampus (44.5%) and hypothalamus (11.6%).

**CONCLUSIONS:** dVRS are always detected in the BG or WM in elderly people, and large dVRS are also prevalent. The topographic distribution of dVRS is not uniform within the brain and may depend on anatomic or pathologic characteristics interacting with aging and sex.

**ABBREVIATIONS:** BG = basal ganglia; 3C = 3 city; CADASIL = cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy; CMA = cortical medullary arteries; dVRS = dilated Virchow-Robin spaces; INSERM = French Institute of Health and Medical Research; LSA = lenticulostriate arteries; UMR = Mixed Unit of Research; WM = white matter

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Virchow-Robin spaces are perivascular spaces that surround small vessels as they extend into the brain tissue. Virchow-Robin spaces can enlarge under undetermined conditions and become dVRS visualized on MR imaging in healthy subjects or in patients with stroke. Evidence is accumulating that dVRS are associated with WM lesions of vascular origin, or dementia, or even multiple sclerosis. Although dVRS are widely detected in healthy individuals or in patients with stroke, their exact prevalence, severity, and topographic distribution in a large population of elderly subjects remains unknown and large epidemiologic studies of this MR imaging marker are still needed.

In previous MR imaging studies of dVRS, major technical limitations were encountered. First, most data were obtained with an MR imaging section thickness larger than 5 mm, though the in-plane resolution could reach 1 × 1 mm² in some studies. Because dVRS are usually of <2 mm in diameter in pathologic data and often present with a linear shape, the limited spatial MR imaging resolution obviously hinders the interpretation of their results. Second, almost all previous studies were based on the assessment of dVRS only in the axial planes, though 3D MR imaging analysis is of paramount importance to disclose the typical course of dVRS along the vascular trunk and for differential diagnosis with lacunar infarction.

In the present study, we analyzed the prevalence, location, and severity of dVRS in a large community-based sample of elderly people. dVRS were identified and analyzed by using high-resolution 3D MR imaging and were graded by using dedicated severity scores both in the BG and within the WM.

**Materials and Methods**

**Subjects**
The 3C Study is a multicenter cohort study, conducted in 3 French cities (Bordeaux, Dijon, and Montpellier) and designed to estimate...
the risk of dementia attributable to vascular risk factors. To be enrolled, subjects had to be ≥65 years of age, noninstitutionalized, and registered on electoral rolls. The detailed description of the study protocol has been previously reported. The study was approved by the ethics committee of the University Hospital of Kremlin-Bicêtre. Each participant signed an informed consent.

The participants from Dijon who were younger than 80 years of age and enrolled between June 1999 and September 2000 (n = 2763) were scheduled to have a cerebral MR imaging examination. Exclusion criteria for the examination were the following: cardiac pacemaker; valvular prosthesis, other internal electrical/magnetic devices; history of neurosurgery/aneurysm; claustrophobia; and the presence of metal fragments (eyes, brain, and spinal cord). Although 2285 subjects (82.7%) agreed to participate, only 1924 subjects underwent brain MR imaging because of financial limitations, among whom 1876 were available for the current analysis.

Individuals with dementia, brain tumors, and self-reported history of stroke were excluded from the current analysis (n = 58), and the final sample size comprised 1818 subjects.

**MR Imaging Data**

MR imaging was performed on a 1.5T Magnetom scanner (Siemens, Erlangen, Germany). A 3D high-resolution T1-weighted brain volume was acquired by using a 3D inversion-recovery fast-spooled gradient-echo sequence (TR = 97 ms; TE = 4 ms; TI = 600 ms; coronal acquisition). The axially reoriented 3D volume matrix size was 256 × 192 × 256 with a 1.0 × 0.98 × 0.98 mm³ voxel size. There were 124 sections covering the whole brain. T2- and proton attenuation–weighted brain volumes were acquired by using a 2D dual spin-echo sequence with 2 TE (TR = 4400 ms; TE1 = 16 ms; TE2 = 98 ms). T2 and proton-attenuation acquisitions consisted of 35 axial sections 3.5-mm thick (0.5-mm between-section spacing) having a 256 × 256 matrix size and a 0.98 × 0.98 mm³ in-plane resolution.

**3D MR Imaging Analysis**

Image analysis was performed on MRicro 1.40 (http://www.sph.sc.edu/comd/rorden/mricro.html), a free image-processing program operating on a Microsoft Windows platform. For each subject’s scan, the first step of MR imaging analysis was performed with T1-weighted images at ×2 magnification on a 27-inch screen to visualize the characteristics of lesions simultaneously in axial, coronal, and sagittal planes. Thus, all suspected lesions were analyzed on the basis of their shape and location in the 3D space (Fig 1). T2- and proton attenuation–weighted images were analyzed in a second step to confirm that the signal intensity of the lesion corresponded to that of CSF.

**Rating of dVRS**

dVRS were defined as CSF-like signal-intensity (hypointense on T1 and hyperintense on T2) lesions that were round, ovoid, or linear; <3 mm in their maximum diameter, with smooth delineated contours; and located in areas supplied by perforating arteries (Fig 2). For lesions fulfilling the same criteria except that their diameter was ≥3 mm, further effort was needed to differentiate them from infarcts by using multiplanar reformatting. Only those with a typical vascular shape and following the orientation of perforating vessels (including cystic lesions with an extension of vascular shape) were then regarded as dVRS (Fig 1).

For each subject, all 124 axially oriented T1-weighted sections were first examined to evaluate the global burden of dVRS and to identify the section containing the largest number of dVRS in both the BG and cerebral WM. In BG, dVRS were then rated according to a 4-level severity score in the section containing the greatest number of dVRS. The degrees of dVRS were defined as follows: degree 1, when there were <5 dVRS; degree 2, when there were between 5 and 10 dVRS; degree 3, when there were >10 dVRS but they were still countable; and degree 4, when an innumerable number of dVRS resulted in a cribriform change in the BG (Fig 3). In cerebral WM, dVRS were scored as follows: degree 1, when there were <10 dVRS in the total WM; degree 2, when there were >10 dVRS in the total WM but <10 in the section containing the greatest number of dVRS; degree 3, when there were between 10 and 20 dVRS in the section containing the greatest number of dVRS; and degree 4, when there were >20 dVRS in the section containing the greatest number of dVRS (Fig 4). This rating scheme was adopted after testing different visual rating methods, including those reported in the literature on a subset of MR imaging data from the first 150 subjects of the cohort and after performing step-by-step multiple rectifications. dVRS with axial diameters of ≥3 mm were defined as “large dVRS” in our study. Their presence, number, and location were systematically recorded.

All images were analyzed by the same experienced reader (Y.-C.Z.) who was blinded to all clinical data. When this reader was uncertain of the categorization of MR imaging signal-intensity abnormalities (n = 113), the corresponding lesions were reviewed by 2 other readers who had extensive experience in MR imaging studies (H.C. and C.T.) and a decision was made on the basis of the majority vote of the 3 readers.

**Other MR Imaging Parameters**

The gray matter, WM, and CSF volumes were estimated with voxel-based morphometry methods detailed elsewhere. Total intracranial volume was computed as the sum of the gray matter, WM, and CSF volumes and brain parenchymal fraction as the ratio of brain tissue volume to total intracranial volume.
Statistical Analysis

Associations between qualitative variables and dVRS scores in the BG or WM were assessed by using \( \chi^2 \) tests. In multivariable analyses, the associations between qualitative variables and dVRS degree in the BG or WM were assessed by using logistic regression controlling for brain size (estimated by total intracranial volume). Associations between continuous variables and dVRS degrees in the BG or WM were examined by using analysis of covariance controlling for total intracranial volume. Potential interactions were tested by adding cross-products into the models. All analyses were performed by using SAS, Version 9.1 (SAS Institute, Cary, North Carolina).

Results

The sample comprised 1818 participants, 61.2% of whom were women. The mean age was 72.5 ± 4.1 years. The intrarater agreement for the rating of dVRS in the BG and WM was...
but not in the WM (men increased with the severity of dVRS in the BG (Table 1). However, a discrepancy of 2 large dVRS (there were 948 large dVRS in total). Large dVRS were observed in the following areas by decreasing frequency: proximal LSA, 19.9%; CMA, 10.2%, hippocampus, 5.8%, distal LSA, 2.8%, brain stem, 1.2%; and hypothalamus, 0.1%. Therefore, half of the large dVRS were detected in the proximal part of LSA, and one-fourth along the CMA. The prevalence of large dVRS in the proximal part of LSA, in the distal part of LSA, or along CMA significantly increased with the severity of dVRS in the BG and WM (P ≤ .02 for all tests).

### Table 1: Cross table of dVRS scores in basal ganglia and white matter

| Scores of dVRS in BG | Degree 1 | Degree 2 | Degree 3 | Degree 4 | Total No. (%) |
|----------------------|----------|----------|----------|----------|---------------|
| Degree 1             | 295 (16.2) | 528 (29.0) | 127 (7.0) | 26 (1.4) | 976 (53.7) |
| Degree 2             | 104 (5.7)  | 344 (18.9) | 147 (8.1) | 46 (2.5) | 641 (35.3) |
| Degree 3             | 26 (1.4)   | 88 (4.8)   | 45 (2.5)  | 18 (1.0) | 177 (9.7)   |
| Degree 4             | 4 (0.2)    | 10 (0.6)   | 6 (0.3)   | 4 (0.2)  | 24 (1.3)    |
| Total                | 429 (23.6) | 970 (53.4) | 325 (17.9) | 94 (5.2) | 1818 (100%) |

### Table 2: Distribution of scores of dVRS in the BG and WM by age and sex

| Location | No. | Mean age (yr) (SD) | P Value | % Men (No.) | P Value |
|----------|-----|--------------------|---------|-------------|---------|
| BG       |     |                    |         |             |         |
| Degree 1 | 976 | 71.7 (3.9)         | <.0001a  | 36.5 (356)  | .0002a  |
| Degree 2 | 641 | 73.0 (4.2)         | 37.8 (242)| 50.3 (88)   | .51c     |
| Degree 3 | 177 | 74.2 (4.0)         | .0001b   | 79.2 (19)   | .51c     |
| Degree 4 | 24  | 75.0 (3.4)         | .51c     |             |         |
| WM       |     |                    |         |             |         |
| Degree 1 | 429 | 72.2 (4.2)         | .004a, .004a | 33.6 (144) | .51b, .53d |
| Degree 2 | 970 | 72.4 (4.1)         | 39.5 (283)| 52.5 (137)  | .51c     |
| Degree 3 | 325 | 72.7 (4.2)         | 42.2 (137)| .51c        |         |
| Degree 4 | 94  | 73.8 (4.1)         | 44.7 (42) |             |         |

* Computed from analysis of covariance controlling for total intracranial volume.
* Computed from analysis of covariance controlling for total intracranial volume and sex.
* Computed from logistic regression controlling for total intracranial volume.
* Computed from logistic regression controlling for total intracranial volume and age.

Discussion

In this sample of 1818 elderly subjects free of dementia or stroke, we found that dVRS were a common MR imaging finding. Using high-resolution 3D MR imaging, we found dVRS in every subject and dVRS of >3 mm in approximately one-third of them. The results also showed that the number of dVRS increased with age in the BG as well as in the WM and that men had more dVRS in the BG than women.

The observation of dVRS in all subjects of this cohort contrasts with the 38% prevalence reported in MR imaging studies of 816 outpatients of wide age range and 3% of 1250 pediatric patients. The different spatial resolutions of imaging may be the main source of this discrepancy. Herein, we used 3D MR imaging acquisitions with a voxel size of 1.0 × 0.98 × 0.98 mm³, whereas most previous studies of dVRS including the above-mentioned were based on 2D MR imaging acquisitions and image sections of 5-mm thickness. Moreover, Groeschel et al found a higher detectability of dVRS on coronal T1-weighted images than on axial T2, sagittal T2, or axial fluid-attenuated inversion recovery images. These data emphasized the importance of section orientation and MR imaging sequences in the assessment of dVRS. In the present study, the 3D T1 imaging analysis used for identification of dVRS may have also increased the detection of dVRS. Virchow-Robin spaces were also visible in 100% of subjects in another study using high-resolution 3D MR imaging in 125 healthy subjects between 1 and 30 years of age, supporting our inferences.

Although a diameter of 3 mm has been widely used as a cutoff value to discriminate silent brain infarcts from dilated perivascular spaces on MR imaging, dVRS with diameters of >3 mm are frequently detected. One-third of individuals in the 3C-Dijon MR imaging sample had 1 or more dVRS larger than 3 mm. This is 3 times higher than the prevalence of lacunar infarctions in the same sample. The most common location of large dVRS was found to be at the proximal part of LSA, in agreement with the most widely recognized location of dilated perivascular spaces. The prevalence of large dVRS was twice as high in this area as along the CMA and 10 times higher than in the distal part of LSA. Various mechanisms underlying the expansion of perivascular spaces have been previously suggested, such as the following: 1) changes in the permeability of the arterial wall, 2) altered drainage and accumulation of amyloid proteins along the vessels, 3) spiral elongation of blood vessels, or cerebral atrophy.
In this study, an accumulation of uncountable dVRS in the BG was observed in 1.3% of participants. A special MR imaging change of cribiform cavities in the BG was the feature of this group. An “état crible” or “status cribrosus” has been previously proved to be multiple dVRS in pathologic studies and has been reported only among cases of vascular Parkinsonism, subcortical ischemic vascular dementia, or CADASIL but has not been reported in the general population, to our knowledge. The present data further emphasize that the accumulation of multiple dVRS is not rare and can occur in elderly people in the absence of severe disability or dementia.

The association between age and the severity of dVRS was consistently detected in previous dVRS studies in hospitalized patients and in patients with lacunar infarction, as well as in specific conditions such as CADASIL. This may suggest 1 or several potential causes of dVRS with aging. In this study, a higher grade of dVRS in the BG was observed more frequently in men than in women. This difference was independent of the total intracranial volume. Most interesting, this sex effect was not detected for dVRS in the WM. Whether variations in ultrastructural characteristics or permeability properties of vessels located within the BG are involved in the male sex association with higher severity of dVRS in the BG will require specific investigation.

In this sample, we also observed that dVRS are commonly detected in areas not usually considered for MR imaging assessment of dilated perivascular spaces. Although still controversial, the hippocampal cavities located between the cornu ammonis and dentate gyres were found at postmortem examination to be covered by a single cell layer, to contain blood vessels, and to be surrounded by intact cerebral parenchyma, which strongly suggest typical dVRS. We found that these cavities are of uniform shape and location, with a frequency of 44.5% in the 3C-Dijon MR imaging cohort, close to the 39% reported by Sasaki et al in 109 patients from 8 to 85 years of age. Whether their severity is related to age and apolipoprotein E4 or E2 alleles as suggested in small samples will need additional investigations. We also found that >10% of patients presented with dVRS in the hypothalamus, a feature never previously reported, to our knowledge.

The strengths of this study include the population-based design, the large sample size, and the use of high-resolution MR imaging and 3D image analysis. The definition of dVRS was also based on strict anatomic and imaging criteria, and the method of assessment proved to be highly reliable. The main limitations include the potential misclassification of dVRS versus small ischemic lesions despite the training of the reader in using 3D imaging analysis and the use of a semiquantitative evaluation for grading of severity.

Conclusions

The results of this study emphasize that large dVRS are highly prevalent in the healthy population and can be detected in most brain structures by using 3D high-resolution MR imaging. The results confirmed that the topographic distribution of dVRS is not uniform within the brain and may depend on anatomic or pathologic characteristics interacting with aging and sex. Whether the severity of dVRS at MR imaging is related to vascular risk factors and/or to other imaging markers of small vessel diseases in healthy elderly people will need additional investigation.

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References

1. Macullich AM, Wardlaw JM, Ferguson KJ, et al. Enlarged perivascular spaces are associated with cognitive function in healthy elderly men. J Neurol Neurosurg Psychiatry 2004;75:1519–23
2. Heier LA, Bauer CJ, Schwartz L, et al. Large Virchow-Robin spaces: MR-clinical correlation. AJNR Am J Neuroradiol 1989;10:209–16
3. Udaka F, Sawada H, Kameyama M. White matter lesions and dementia: MRI-pathological correlation. Ann N Y Acad Sci 2002;977:411–15
4. Patankar TF, Mitra D, Varma A, et al. Dilation of the Virchow-Robin space is a sensitive indicator of cerebral microvascular disease: study in elderly patients with dementia. AJNR Am J Neuroradiol 2005;26:1512–20
5. Wuerfel J, Haertle M, Waiczies H, et al. Perivascular spaces: MRI marker of inflammatory activity in the brain? Brain 2008;131:2332–40
6. Bokura H, Kobayashi S, Yamaguchi S. Distinguishing silent lacunar infarction from enlarged Virchow-Robin spaces: a magnetic resonance imaging and pathological study. J Neurol 1999;245:116–22
7. Grooschel S, Chong WK, Suttees R, et al. Virchow-Robin spaces on magnetic resonance images: normative data, their dilatation, and a review of the literature. Neuroradiology 2006;48:745–54
8. Alperovich A. Vascular factors and risk of dementia: the design of the three-city study and baseline characteristics of the study population. Neuroepidemiology 2003;23:316–25
9. Soumare A, Elbaz A, Zhu Y, et al. White matter volume and motor performances in the elderly. Ann Neurol 2009;65:706–15
10. Lemaitre R, Crivello F, Grissiat B, et al. Age- and sex-related effects on the neuroanatomy of healthy elderly. Neuroimage 2005;26:900–11
11. Rollins NK, Deline C, Morris MC. Prevalence and clinical significance of dilated Virchow-Robin spaces in childhood. Radiology 1993;189:53–57
12. Rouhl RP, van Oostenbrugge RJ, Knottnerus II, et al. Virchow-Robin spaces relate to cerebral small vessel disease severity. J Neurol 2008;255:692–96
13. Vermeer SE, Den HT, Koudstaal PJ, et al. Incidence and risk factors of silent brain infarcts in the population-based Rotterdam Scan Study. Stroke 2003;34:392–96
14. Takao M, Koto A, Tanahashi N, et al. Pathologic findings of silent, small hypointense foci in the basal ganglia and thalamus on MRI. AJNR Am J Neuroradiol 2003;24:3114–18
15. Pollock H, Hutchings M, Weller RO, et al. Perivascular spaces in the basal ganglia of the human brain: their relationship to lacunes. J Anat 1999;195(Pt 2):761–74
16. Mancardi GL. Neuropathologic study of lacunae and cribiform cavities of the brain. Eur Neurol 1989;29(4):16–19
17. Penelon G, Gray F, Wallys C, et al. Parkinsonism and dilatation of the perivascular spaces (état criblé) of the striatum: a clinical, magnetic resonance imaging, and pathological study. Mov Disord 1995;10:754–60
18. Braffman BH, Zimmerman RA, Trojanowski JQ, et al. Brain MR: pathological correlation with gross and histopathology. 1. Lacunar infarction and Virchow-Robin spaces. AJR Am J Roentgenol 1988;151:551–58
19. Roman GC. On the history of lacunes, état criblé, and the white matter lesions of vascular dementia. Cerebrovasc Dis 2002;13(5):264–72
20. Cumurciuc R, Guichard JP, Reizine D, et al. Dilation of Virchow-Robin spaces in CADASIL. Eur J Neurol 2006;13:187–90
21. Sasaki M, Sone M, Ehara S, et al. Hippocampal sulcus remnant: potential cause of change in signal intensity in the hippocampus. Radiology 1993;188:743–46
22. Bastos-Leite AJ, van Waesberghe JH, Oen AL, et al. Hippocampal sulcal cavities on MRI: relationship to age and apolipoprotein E genotype. Neurology 2000;54:2150–53