Giant Tumefactive Perivascular Spaces

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BACKGROUND AND PURPOSE: The brain perivascular spaces (PVSs) are pial-lined, interstitial fluid-filled structures that accompany penetrating arteries. When enlarged, they may cause mass effect and can be mistaken for more ominous pathologic processes. The purpose of this study was to delineate the broad clinical and imaging spectrum of this unusual condition.

METHODS: Thirty-seven cases of giant PVSs were identified from 1988 to 2004 and were retrospectively reviewed. Clinical data collected included patient demographics, presenting symptoms, and follow-up. Histopathologic data were reviewed when available. Images were evaluated for size and location of the giant PVSs, associated mass effect, hydrocephalus, adjacent white matter changes, and contrast enhancement.

RESULTS: There were 24 men and 13 women with an age range of 6–86 years, (mean 46 years). The most common presenting feature was headache (15 patients). Thirty-two cases had multilocular clusters of variably sized cysts. Five lesions were unilocular. All lesions had signal intensity comparable to CSF and did not enhance. The most common location for the giant PVSs was the mesencephalothalamic region (21/36). Fourteen were located in the cerebral white matter; two were in the dentate nuclei. Nine giant mesencephalothalamic PVSs had associated hydrocephalus, which required surgical intervention.

CONCLUSION: Giant tumefactive PVSs most often appear as clusters of variably sized cysts that are isointense relative to CSF and do not enhance. They are most common in the mesencephalothalamic region and may cause hydrocephalus. Although they may have striking mass effect, giant PVSs should not be mistaken for neoplasm or other diseases.

The perivascular spaces (PVSs) of the brain, also known as Virchow-Robin spaces, are pial-lined interstitial fluid (ISF)-filled structures that accompany penetrating arteries and arterioles for a variable distance as they descend into the cerebral substance (1–3). Recent studies have shown that the PVSs are surprisingly complex entities with significant variability in both ultrastructure and possible function. Routinely, PVSs in many areas of the brain can be identified on MR images obtained in patients of all ages (4). Occasionally the PVSs may become strikingly enlarged, causing mass effect and assuming bizarre configurations that may be mistaken for a more ominous disease, such as a cystic neoplasm (Figure 1). While scattered cases of giant tumefactive PVSs have been reported (5–7), no large series of these lesions has been presented. The purpose of this study is to delineate the broad clinical and imaging spectrum of giant tumefactive perivascular spaces to further characterize this unusual lesion.

Methods

We performed a retrospective review of all cases of giant PVSs referred for imaging consultation or treated at our institution over a 16-year period spanning 1988 to 2004. Most cases referred for second opinion had the presumptive diagnosis of cystic neoplasm.

We defined giant PVSs as equal to or greater than 1.5 cm, occurring as either a solitary lesion or in clusters of multiple contiguous cysts.

Clinical data collected included patient age, sex, and presenting symptoms. Histopathologic data in patients who underwent biopsy were collected and reviewed. The specific histologic stains used included: Hematoxylin and Eosin as a general histologic stain; Hematoxylin van Gieson to stain the collagen of the meninges on the outer aspect of the cyst wall; and reticulin to stain the collagen of the meninges on the outer aspect of the cyst wall.
All patients were imaged with MR imaging using a variety of 1.5T MR imaging units. Typical pulse sequences included T1-weighted spin-echo images, fast spin-echo T2-weighted images, proton density–weighted (PD) images, and fluid attenuated inversion recovery (FLAIR) images. Patients’ MR imaging examinations included either a PD or FLAIR sequence. Gadolinium-enhanced T1-weighted imaging was also performed in most patients (34/36). Two studies were performed without the use of contrast medium. Ten patients underwent CT performed with 5-mm axial sections.

Images were evaluated to determine the size of the giant PVs, establish whether cysts were unilateral or bilateral, and ascertain whether cysts were solitary or clustered. Images were also assessed for the appearance of associated focal or generalized mass effect, hydrocephalus, adjacent white matter changes, and presence of contrast enhancement.

Results

Demographics

Thirty-seven patients with giant PVs were identified, including 24 men and 13 women (M:F = 1.8:1). Patient age ranged from 6 to 86 years with a mean of 46 years. Five unilocular lesions were identified ranging in size from 1.5 cm to 3.5 cm. Multiple clustered cysts (32/37 cases) were present in most cases. The clusters were all greater than or equal to 1.5 cm in size, ranging from 1.5 cm to 10 cm in greatest dimension. There were 27 unilateral lesions and 10 bilateral lesions.

Clinical Findings:

Complete clinical data were available in 33 patients with a partial history available in the remaining four patients. The most common presenting symptom was headache, present in 15/33 patients. Other presenting features included dizziness, dementia, visual changes, post-traumatic evaluation, seizure, syncope, stroke, memory problems, cranial neuropathy and poor balance and concentration. One additional patient had a 6th nerve palsy which was found to be related to diabetic neuropathy and resolved on follow-up imaging. One patient who presented with visual changes was ultimately diagnosed with uveitis, which was treated and resolved (Table 1).

Imaging Findings

CT was performed in 10 of 37 patients. All giant PVs were low-attenuation lesions, isoattenuated relative to CSF, which showed no enhancement following contrast medium administration. No calcifications or other associated abnormalities were identified.

All giant PVs were isointense relative to CSF signal intensity on all MR images regardless of pulse sequence. Contrast-enhanced images were available in 35/37 cases and showed no enhancement. Diffusion-weighted (DW) imaging was available in eight of 37 cases and showed no diffusion restriction.

Lesions were subdivided by location (Table 2). There were 21 patients with lesions involving the midbrain or thalamus (mesencephalothalamic area) (Fig 2) and 14 lesions predominately involved the hemispheric or subcortical cerebral white matter. Two lesions were present in the region of the dentate nuclei of the cerebellar hemispheres.

Of the 21 mesencephalothalamic lesions, four were bilateral. No signal intensity alteration was present in the adjacent brain parenchyma. Nine of these 21 mesencephalothalamic lesions had associated hydrocephalus and were treated with ventriculostomy (5), ventriculoperitoneal shunting (3), or cystoperitoneal shunting (1) (Fig 3).

There were 14 patients with giant PVs in the cerebral white matter (Fig 4). Five of these white

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N = 33
matter cases were bilateral. White matter lesions were located in the subcortical white matter (8/14), hemispheric white matter (4/14), or both (2/14). Most white matter lesions involved the frontal lobe (8/14) with other areas of involvement including the parietal lobe (3/14), temporal lobe (2/14), and occipital lobe (1/14). There was involvement of the cingulate gyrus subcortical white matter in four cases and three cases had involvement of the corpus callosum. In these cases, the gray matter was stretched over the giant PVSs (Fig 5).

Associated white matter signal intensity alterations occurred in seven of the cerebral white matter cases. Two of these were focal changes, with increased T2 and FLAIR signal intensity, surrounding the giant PVSs, and were found in older patients (a 68-year-old woman and a 73-year-old man) (Fig 4). Five additional patients with lesions in the white matter had more diffuse, confluent abnormal signal intensity alteration consisting of increased T2 and FLAIR signal intensity surrounding the PVSs (Fig 6).

There were two lesions that involved the region of the cerebellar dentate nuclei. In one case, the lesions were bilateral. There was no associated signal intensity alteration in the adjacent brain parenchyma (Fig 7).

**Surgical and Pathologic Findings**

Surgery was performed in 12 patients. In seven cases, biopsy was performed. Histopathologic results in most cases (6/7) disclosed a pial-lined cyst with no evidence of neoplasm or infection. One patient with a midbrain lesion had a layer of pia and arachnoid on the outer aspect of the tissue but no pial lining. This patient had a small fragment, approximately 1 mm cubed, of the cyst lining sent to the pathology department. It was fixed in formalin, embedded in paraffin, and stained with hematoxylin and cosin, hematoxylin van Gieson for collagen, and by immunocytochemistry for glial fibrillary acidic protein (GFAP) for astrocytes Figure 8. Figure 8A shows the cyst lining consists of a very thin layer of gliotic brain tissue coated on its outer aspects by collagenous material related to the pia arachnoid on the surface of the midbrain. No lining of pia matter was identified on the inner cyst surface. The brain tissue lining the cyst was extensively gliotic as shown in the GFAP preparation (Fig 8B). Close examination of the cells in the brain tissue lining the cyst shows that it is mainly composed of astrocytes (Fig 8C). The absence of a lining of pia matter is only to be expected in this case, as the cyst was very large and it is unlikely that the thin delicate lining surrounding a vessel would be stretched to this extent. When there is less extensive enlargement of perivascular spaces, the lining of pia matter may remain. Two patients had a ventriculoperitoneal shunt placed, and the size of the giant PVSs was stable in each case. Another patient had a cystoperitoneal shunt placed, and there was a decrease in the size of the giant PVS from 10 cm to 5 cm. In this case, the patient's headache also improved. However, the fluid reaccumulated after the patient had a minor traumatic insult, and the giant PVS enlarged to near-presentation size. One patient underwent PVS drainage followed by ventriculoperitoneal shunt placement after the PVS returned to its original size. Five patients underwent a third ventriculostomy; three of these patients also had a biopsy of the cyst wall.

Follow-up MR imaging was available in seventeen patients from 1–12 years after the initial diagnosis. No interim change was seen on repeat MR imaging in any case.

**Discussion**

When PVSs become markedly expanded, they can be misinterpreted as other pathologic processes, most often a cystic neoplasm. As most of these cysts border a ventricle or subarachnoid space, reports of such cases have offered an extensive differential diagnosis that includes cystic neoplasms, parasitic cysts, ventricular diverticula, cystic infarction, non-neoplastic neuropathelial cysts, and deposition disorders such as mucopolysaccharidosis (4, 7–9). The aim of this study...
was to delineate the classic imaging appearance of giant PVSs in an effort to prevent misdiagnosis and unnecessary biopsy.

Typical PVSs occur in many locations. The most common site is along the lenticulostriate arteries, just above the anterior perforated substance and adjacent to the anterior commissure. Less commonly, PVSs occur along arteries that have penetrated the cerebral cortex and extended into the white matter (4, 8). Other areas where prominent PVSs can be identified include the subinsular region, dentate nuclei, and cerebellum (2). PVSs have typical MR imaging features as previously described. They are round or oval with a well-defined, smooth margin, occur along the path of penetrating arteries, are isointense relative to CSF, and demonstrate no enhancement following contrast medium administration (4, 6, 8, 9). When PVSs become enlarged, they are known as giant PVSs, “cavernous dilatation,” or Poirier’s Type IIIb “expanding lacunae” (6, 10, 11).

Giant PVSs are expanded PVSs that occur along the penetrating vessels, most commonly in the mesencephalothalamic region in the territory of the paramedial mesencephalothalamic artery and in the cerebral white matter (4, 7, 10). They differ from typical PVSs in that they are larger in size and may have associated focal mass effect. In addition, white matter giant PVSs may have associated T2 and FLAIR signal intensity alteration in the adjacent white matter.

Patients usually present with nonspecific findings that are not attributable to the giant PVSs. In our series, headache was the most common presenting feature and occurred in approximately 50% of our patients. Other presenting complaints included dizziness, dementia, visual changes, post-trauma, seizure, syncope, memory problems, poor balance, and poor concentration. Several studies have found that the clinical features of patients with this type of lesion do not correlate with the imaging findings or have been found incidentally at autopsy (9,12–14). However, widespread dilatation of the PVSs has been reported in association with dementia and parkinsonism (15, 16). Some authors have correlated large convexity and white matter PVSs with increasing age (4). Others have concluded that the prevalence of basal ganglia PVSs in patients younger than 40 years is not significantly different from that of older patients (8). We have observed prominent PVSs on high-spatial-resolution MR images in patients of all ages and in all typical anatomical locations. The clinical symptoms did not correlate with the giant PVSs in our series, except in those cases that had associated hydrocephalus, requiring CSF diversion.

Imaging of enlarged PVSs is characteristic (17, 18). All cases have associated mass effect, from mild to severe. The mass effect may occasionally cause ob-
constructive hydrocephalus as was seen in eight of our study patients, all with lesions in the mesencephalo-thalamic region. Associated hydrocephalus has been reported in the literature and may require surgical

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**Fig 4.** Sagittal T1-weighted (A), axial FLAIR (B), and coronal T2-weighted (C) images obtained in a 71-year-old man with dementia show extensive involvement of the hemispheric and subcortical white matter with multilocular giant perivascular spaces. The coronal image shows the marked asymmetry of the lesions. Note the scattered focal white matter changes surrounding some of the lesions (black arrows) seen best on the FLAIR image. Case courtesy of Anthony Doyle, MD.

**Fig 5.** Sagittal T1-weighted (A), axial FLAIR (B), and axial T2-weighted (C) images obtained in a 46-year-old woman with a visual field defect show extensive involvement of the corpus callosum and cingulate gyrus (A, white arrow) with extension to the subcortical white matter of the parietal and occipital lobes. There is slight increased signal intensity surrounding the lesions, best seen on FLAIR image (B, white arrows). The gray matter is stretched and displaced over the multiloculated giant perivascular spaces. Case courtesy of Leena Valanne, MD.

**Fig 6.** Images obtained in a 37-year-old man with headaches. Axial T2-weighted (A) and axial FLAIR (B) images show diffuse, confluent white matter hyperintensity surrounding the giant perivascular spaces. There is mild gyral expansion over the perivascular spaces. The diffusion-weighted image (C) shows no diffusion restriction.
shunt surgery (19). FLAIR imaging typically shows complete signal intensity suppression without abnormalities in the adjacent parenchyma. In a few reported cases, small foci of high signal intensity adjacent to the cystic spaces have been identified (20). Our study shows that giant PVSs that occur in the white matter may have surrounding signal intensity abnormality seen on T2-weighted or FLAIR images, as seen in half of our cases. Lesions with adjacent white matter changes can be divided into two groups. In elderly patients in our series, the white matter changes surrounding the dilated PVSs were discrete areas of abnormal T2 and FLAIR hyperintense signal intensity abnormality. One possible theory is that this associated signal intensity alteration may represent advanced chronic ischemic change related to mass effect of the PVSs (5). Another possibility is that dilated PVSs may result from chronic mechanical stress caused by high blood pressure on the brain arterioles (21). However, most hypertensive patients imaged do not have associated giant PVSs.

In younger patients in our series, the signal intensity abnormality was more diffuse, with confluent T2 and FLAIR increased signal intensity surrounding the giant PVSs. Although the exact etiology is not known, we suggest that the changes may be secondary to gliosis or spongiosis. Alternatively, the signal intensity changes may be related to multiple tiny tightly clustered PVSs that are too small to be discriminated on the basis of current MR imaging findings. Abnormal hyperintensity in the cerebral white matter on T2-weighted images occurs in more than 30% of the neurologically healthy elderly population (9, 22, 23). Histopathologic study with MR imaging correlation has indicated that many of these lesions are PVSs (8, 24–26). One author found that PVSs greater than 2 mm were found in 67 (8%) of 816 patients (4).

Giant PVSs (up to 2–3 cm in diameter) have been reported to occur as a normal variant (4, 27). The precise etiology of these enlarged PVSs is unknown. Spiral elongation of the penetrating blood vessels has been suggested as one possible cause of this phenomenon (24). Increased CSF pulsations, the ex vacuo phenomenon, or an abnormality of arterial wall permeability have also been cited as contributing factors (9, 13, 28, 29).

Experimental studies suggest that the periarterial spaces are the routes by which interstitial fluid (ISF) drains from brain tissue (30). In humans, amyloid-beta (Aβ) appears to be entrapped within the drainage pathways in the cerebral amyloid angiopathy that accompanies Alzheimer disease (AD) (31, 32). Although the periarterial spaces in the cortex are not dilated in AD, periarterial spaces in the white matter are expanded possibly due to obstruction of ISF drainage pathways by Aβ in the proximal parts of each artery (33). By analogy, obstruction of ISF drainage pathways could be a major factor in the pathogenesis of giant PVSs, although the exact nature of the obstruction is unclear.

Other investigators have postulated that because the pial membrane surrounding the penetrating ar-
tery is fenestrated, accumulation of brain interstitial fluid between the vessel and pia or within the inter-pial space causes the PVSs to enlarge (27). We found that giant PVSs are most common in the mesencephalothalamic region, which may indeed be related to the two layers of pia in this location. In addition, the significant mass effect with associated obstructive hydrocephalus seen in the mesencephalothalamic group may, in part, be related to the two layers of pia in this location. In addition, the significant mass effect with associated obstructive hydrocephalus seen in the mesencephalothalamic group may, in part, be related to the two pial layers.

Giant PVSs with mass effect may occur as single or multiple clustered cysts that can be mistaken for more ominous disease (34). When the lesions in question occur in a characteristic location along the path of a penetrating vessel, follow CSF signal intensity on all sequences, do not enhance with contrast material, and have normal adjacent brain parenchyma, their appearance is virtually pathognomonic of giant PVSs. An extensive differential diagnosis is superfluous and biopsy unnecessary (14).

Conclusion

Giant tumefactive PVSs are interstitial-fluid filled structures that accompany arteries and arterioles as they penetrate the brain. They have characteristic imaging features: round or oval, single or multilocular lesions that are isointense relative to CSF regardless of imaging sequences and do not enhance. They are most common in the mesencephalothalamic region and may have associated obstructive hydrocephalus that may require surgical intervention. Although they
References