Enlarged Perivascular Space and Its Correlation with Polysomnography Indicators of Obstructive Sleep Apnea

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Purpose: There is increasing evidence of a causal interaction between obstructive sleep apnea (OSA) and white matter hyperintensity (WMH). WMH and enlarged perivascular space (EPVS) are the neuroimaging markers for cerebral small vessel disease (CSVD). Thus, this study aimed to determine whether a contextual relationship existed between OSA and EPVS.

Participants and Methods: In this study, 107 participants underwent 1-night polysomnography, brain magnetic resonance imaging (MRI) and health screening examinations and were classified as 63 OSA patients (mild, moderate, and severe groups), and 44 healthy controls. We assessed the sleep characteristics in OSA group, quantified the total EPVS from MRI and related them to the measures of polysomnography-obtained sleep parameters.

Results: Polysomnography revealed that 63 OSA patients had sleep architecture alteration. A higher proportion of N2 phase sleep (N2%), lower percentage of N3 sleep (N3%) and REM sleep (REM%), as well as increased arousal index (AI), oxygen desaturation index (ODI) and decreased lowest arterial oxygen saturation (LSaO2) were detected. The results also indicated a higher prevalence and a larger number of EPVS, and a lower Mini Mental State Scale (MMSE) scale score in OSA group. LSaO2, N3% and REM% were negatively correlated with the total EPVS, whereas ODI, AI and N2% were positively correlated with the total EPVS.

Conclusion: The findings suggested that OSA patients had sleep disturbances with a higher incidence and more severe EPVS. Furthermore, the EPVS in OSA might be secondary to sleep disturbances, intermittent hypoxemia and the respiratory event-related hemodynamic changes. Thus, our findings highlighted that increased risk for EPVS in OSA is a potential contributor to increased stroke risk in OSA.

Keywords: obstructive sleep apnea, polysomnography, sleep, mini mental state scale, perivascular space

Introduction

Perivascular space (PVS) is an interstitial fluid-filled channel surrounding smaller arteries and veins in the brain, which plays a role in the proto-lymphatic system, exerts a certain effect on interstitial elimination, and is called the Glymphatic system. The glymphatic system that participates in cerebrospinal fluid (CSF) will re-circulate via the brain, interchange with interstitial fluid (ISF) and remove the protein waste from the brain, including Amyloid-β, tau, and α-synuclein. Glymphatic clearance is temporally and cyclically regulated; in this regard, the
fluid transport is enabled by sleep and suppressed during wakefulness, thereby modulating the balance between protein clearance and aggregation.3 A study indicates that the sleep efficiency is negatively correlated with enlarged PVS (EPVS).4 However, the precise mechanism and frequency of EPVS is still a source of controversy. EPVS has been previously suggested to be associated with normal aging and pathological conditions, such as hypertension and dementia.5 It has been previously suggested that EPVS is related to dementia and stroke.6,7 The EPVS burden is related to cognitive impairment,4,8 which is essential to identify the treatable or preventable causes.

Obstructive sleep apnoea (OSA), a sleep disorder characterized by repeated airway obstruction, leads to hypoxaemia and sleep disturbance.9 As suggested by Song and colleagues, OSA also promoted the occurrence of hypertension and endothelial dysfunction to increase the risk of cerebral small vessel disease (CSVD).10 Previous studies have highlighted that patients affected by OSA suffer from stroke and dementia.11–15 OSA has also been indicated to precede the development of cognitive dysfunction, dementia and stroke within the community-based elderly.16,17

OSA is related to abnormalities in structures, functions and metabolic brain imaging, and its characteristics are compatible with damage in neurons.18–21 Besides, white matter hyperintensity (WMH) and EPVS have become one of the recognized imaging markers for CSVD. However, the majority of prior studies have focused on OSA and WMH or silent cerebrovascular lesion cases.19,22 Some articles have investigated EPVS in OSA, but there are no consistent results regarding how OSA affects EPVS.20 Del et al documented that nocturnal desaturation and moderate-severe OSA were related to WMH or PVS,11,22 but some literature proposes inconsistent opinions.23 These studies report individuals who are older, so these samples cannot represent the general population. Thus, whether OSA serves as a risk factor for EPVS among the young and middle-aged populations remains unclear.

This work was carried out aiming to observe the changes in PVS in patients with OSA by brain magnetic resonance imaging (MRI), explore the possible mechanism of PVS expansion in patients with OSA, and examine the associations between these two conditions and the clinical value of early detection and treatment of OSA in evaluating the risk of stroke caused by OSA.

Participants and Methods
Study Sample
From January 2018 to December 2019, 200 physical examinees (including 108 men and 92 women) aged 18–60 years admitted for MRI and overnight polysomnography (PSG) at the sleep unit of the Second Affiliated Hospital of Zhengzhou University were enrolled for the present analysis. During the night of PSG, participants were under the supervision of a qualified technician and underwent PSG monitoring for at least 7 h. All subjects provided informed consent for retrospective analyses of their data. The study protocol was approved by the Second Affiliated Hospital of Zhengzhou University (NO. 2021132). This work was carried out according to the Declaration of Helsinki.

The inclusion criteria included the OSA group that received PSG monitoring at our hospital and conformed to the related standards:24 the apnea hypopnea index (AHI) ≥5/h; and one or more sleep-related symptoms (snoring, witnessed apneas, daytime sleepiness, and non-restorative sleep). Controls were included according to the criteria of AHI<5/h and no medical or psychiatric disease.

The exclusion criteria were as follows: those received less than 9 years of education; the diagnosis of dementia; anxious and depressive disorder; unable to cooperate with the investigation due to hearing, visual and limb activity disorders; neurological disease; psychosis; previous head injury or stroke; sleep disorders in addition to OSA; vascular risk factors (hypertension, diabetes mellitus, dyslipidemia); cardiovascular disease; use of psychoactive medications; structural brain abnormalities; and history of drinking and smoking. The flowchart of subject recruitment is shown in Figure 1.

Polysomnography (PSG)
PSG was conducted using an electrooculogram, a chin and bilateral anterior tibial surface electromyogram, an electrocardiogram, six electroencephalography channels (F3-A1, F4-A2, C3-A1, C4-A2, O1-A1, and O2-A2), nasal and oral airflow sensors, thoracic and abdominal movements sensors, and a Nonin finger probe (Nonin Medical Inc., Plymouth, MN, USA) was utilized for blood oxygen saturation.25 Monitoring was conducted with the Grael-v2 PSG amplifier (Compumedics, Abbotsford, Australia), and the montage electrodes were used for basic sleep diagnosis. Sleep and related events were analyzed according to the American Academy of
Sleep Medicine (AASM) Scoring Manual rules (version 2.4). All sleep data were analyzed by the same physician trained in PSG technology, and related parameters were generated with the Pro Fusion PSG 4 software (Compumedics).

MRI Protocol and Assessment
The Signa HD MRI 3.0 T superconducting magnetic resonance imaging device (GE, USA) was employed for head MRI examinations in all the enrolled subjects. Typically, the specific head coil was utilized for routine axial, sagittal and coronal scans, including the T1WI, T2WI and FLAIR sequences. Of them, T1WI sequence adopted the spin echo sequence (repetition time (TR) = 400ms; echo time (TE) = 8.1ms), T2WI sequence (TR=4080ms; TE=109ms), FLAIR sequence (TR=8600ms; TE=123ms). In addition, the layer thickness was 5 mm and the layer distance was 1.5 mm. During the MRI examination, the subjects were in a supine position on the examining table, kept quiet and awake, closed their eyes, and kept a quiet breathing state, with their hands being placed on both sides of the body.

EPVS Standards
The PVS standards determined by Cumurciuc et al were adopted: (i) based on the different image sections, PVS was round, oval or linear, with clear and smooth boundary, and the diameter was generally < 3mm; (ii) it had consistent direction with the penetrating blood vessel; (iii) PVS had the same signal as cerebrospinal fluid (CSF) on T1WI, T2WI and FLAIR sequences; (iv) there was no enhancement effect or space-occupying effect.

In this study, MRI-visible PVS was defined as EPVS, and the maximum diameter was≤3 mm. EPVS was selected at the locations shown below: (i) frontoparietal subcortical white matter; (ii) centrum semiovale; and (iii) basal ganglia (Figure 2). In each region, we analyzed the left and the right sides, and adopted the greatest score on each side in subsequent statistical analyses. Afterwards, we determined the global score by adding all results acquired from three regions. The examined that plane and EPVS numbers were determined and recorded under the guidance of two experienced MRI neurologists who were blinded to OSA/control status. Any disagreement...
was discussed by the two neurologists, and the examined plane was eliminated when no consensus was reached between them.

**Questionnaire Evaluation**

Mini-Mental State Examination (MMSE) was carried out to assess cognitive function, and the Epworth Sleepiness Scale (ESS) was adopted to estimate the excessive daytime sleepiness. The scale scores were independently completed by the subjects in a quiet environment, and the subjects were asked to sit upright and lean forward slightly between 2:30 pm and 5:30 pm every day, so as to keep the brain alert.

The scale of MMSE ranged from 0 to 30 points, MMSE scores ≥27 revealed a normal cognition, those of 21–26 indicated mild cognitive impairment, those of 11–20 reported moderate cognitive impairment, and those <10 were classified as severe cognitive impairment. The ESS was an eight-item assessment of somnolence, with possible scores ranging from 0 to 24 points. A score of over 10 suggested excessive daytime sleepiness.

**Statistical Methods**

All data were analyzed using the SPSS 20.0 statistical software. For counting data, group comparisons were performed using Chi-square tests. Normality of the measurement data was tested with Shapiro–Wilk test. Normal distributions were described by means ± standard deviations (SD), while abnormally distributed measurement data were expressed as median (quartile) (M [Q25, Q75]). The Kruskal–Wallis test was conducted in group comparisons, and Fisher’s least significant difference (LSD) method was used to adjust for multiple comparisons. Due to the abnormal distribution of data, a nonparametric partial Spearman’s rank-order correlation was utilized to compare the correlations of objective sleep parameters with EPVS. Age, body mass index (BMI) and ESS were incorporated as covariates in the analysis because of their potential influences on both sleep and PVS. Statistical significance was set at p < 0.05.

**Results**

**Sample Characteristics**

Among the 200 subjects enrolled in the present work, two were excluded because of the low MRI scan quality, whereas 61 were ruled out from this work due to diabetes, hypertension, cardiovascular disease or neurological disorder. In the diagnosis process of PSG, six subjects with the overall sleep time <4 h were eliminated from this work. In addition, 23 smokers or drinkers and another one meeting the criteria of current major depression were also eliminated. Therefore, finally, 107 subjects (including 44 (41.12%) females) were enrolled, with the age of (31.72±11.86) years.

Table 1 shows the features of participants with mild, moderate or severe OSA and the control subjects. According to the American Academy of Sleep Medicine guidelines, 44 (41.12%) subjects showing the AHI<5/h were enrolled as the control group; 25 (23.36%) had AHI of 5–15/h (mild); 20 (18.69%) had AHI of 15–30/h (moderate); and 18 (16.82%) had AHI >30/h (severe). Differences in clinical or demographic features were not significant among the different groups, except for BMI and ESS. Nonetheless, relative to control, the MMSE score of OSA group was lower, and that of severe OSA group was significantly lower.

**PSG Parameters Findings**

The inter-group and intra-group PSG results were statistically analyzed (Table 2). Compared with control group, the lowest arterial oxygen saturation (LSaO2), sleep efficiency (SE), sleep latency (SL) and N3%, REM% in OSA group decreased, and the differences were of statistical significance (Z=8.73, p <0.01; Z=3.01, p <0.01; Z=4.25, p <0.01; Z=8.51, p <0.01; Z=7.87, p <0.01). The AHI,
Table 1 Demographic Features and EPVS in Control and OSA Groups

| Parameter          | Control Group (n=44) | Mild Group (n=25) | Moderate Group (n=20) | Severe Group (n=18) | P    |
|--------------------|----------------------|-------------------|----------------------|---------------------|------|
| Sex (male/female)  | 24/20                | 15/10             | 12/8                 | 12/6                | 0.45 |
| Age (year)         | 33.09±13.39          | 30.92±12.39       | 31.50±9.02           | 29.50±9.21          | 0.70 |
| BMI (kg/m²)        | 23.82±3.44           | 24.99±2.41        | 26.38±1.40           | 27.96±2.38          | 0.00 |
| Neck perimeter (cm)| 36.62±1.85           | 36.81±2.14        | 37.41±1.77           | 37.13±2.20          | 0.18 |
| Snoring, n (%)     | 16(36.36%)           | 9(36.00%)         | 8(40.00%)            | 8(44.44%)           | 0.72 |
| ESS (score)        | 3.96±2.35            | 3.92±2.07         | 5.54±2.01            | 6.13±1.27           | 0.02 |
| MMSE (score)       | 28.00±1.29           | 27.84±1.05        | 27.64±1.52           | 27.12±1.58          | 0.11 |
| EPVS, n (%)        | 18(40.91)            | 16(64.00)         | 15(75.00)            | 17(94.44)           | <0.01|
| EPVS (nr)          | 7.14±9.65            | 12.16±10.59       | 26.64±15.98          | 47.13±17.52         | <0.01|

Notes: P was the p-value produced when comparing OSA group with control group; the neck perimeter (in cm) was measured at the level of the cricothyroid membrane. Differences of P<0.05 indicated statistical significance.

Abbreviations: BMI, body mass index; MMSE, Mini Mental State Scale; ESS, Epworth Sleepiness Scale; EPVS, enlarged perivascular space; nr, number.

ODI, wake after sleep onset (WASO), arousal index (AI), N1% and N2% in OSA group increased compared with those in control group, and the differences were statistically significant (Z=8.63, p<0.01; Z=8.52, p<0.01; Z=14.71, p<0.01; Z=3.21, p<0.01; Z=6.57, p<0.01; Z=7.41, p<0.01). Besides, when compared among the three OSA groups, the N1%, N2% and AI in severe group apparently increased relative to those in mild group (all p-values <0.05); in severe group, the LSaO2, N3% and REM% were notably lower than those in mild group (all p-values <0.01).

The Prevalence and Number of EPVS
EPVS was found in 18 (40.91%) control subjects, 16 (64.00%) mild, 15 (75.00%) moderate and 17 (94.44%) severe OSA patients, respectively. Compared with control group, the incidence rate of EPVS in OSA patients was higher (χ²=13.64, P<0.01), and that in severe OSA patients was higher than those in mild and moderate OSA patients (χ²=5.44, P=0.02; χ²=2.69, P=0.10). There were 7.14±9.65, 12.16±10.59, 26.64±15.98 and 47.13±17.52 EPVS in control subjects, mild, moderate and severe OSA groups, separately. The number of EPVS in OSA group was higher than that in control subjects (Z=5.07, P<0.01) (Figure 3), and that in severe group was the highest (Z=3.04, P<0.01; Z=4.96, P<0.01) (Table 1).

Correlations of MMSE, Respiratory Parameters and Sleep Structure with EPVS
The correlations of EPVS with diverse indexes are shown in Tables 3 and 4. After adjusting for age, BMI and ESS, correlations of EPVS numbers with N1% and MMSE were not significant. AHI, ODI, TS90%, AI and N2% were positively correlated with EPVS, whereas LSaO2, SE, N3% and REM% were negatively correlated with EPVS.

Discussion
This study suggested that OSA mainly resulted in sleep structure disturbance, sleep fragmentation and intermittent hypoxemia. The main findings of our study provided evidence for the relationship between OSA and PVS. The prevalence and number of EPVS increased in OSA groups, and such variations were related to OSA severity via the pathophysiologic mechanisms, resulting in cerebrovascular diseases and cognitive decline.

PSG serves as the gold standard to diagnose OSA, and it reflects the objective sleep quality of OSA cases. OSA cases have poor nighttime sleep quality, which has been well recognized among scholars which supports the results in our study. The most fundamental incentive for the brain to sleep lies in its own self-preservation: only the sleeping brain is capable of efficiently clearing the waste products generated during active wakefulness through the perivascular drainage system. The decreased sleep quality will destroy neurotoxin elimination, and disturb brain interstitial fluid drainage and local inflammation. This process may be expected to progressively suppress lymphatic flow in the most affected regions that may result in PVS expansion. Different observations are reported in prior works regarding cerebral PVS in OSA. In this study, we analyzed 63 OSA cases and discovered that the OSA group showed markedly increased EPVS compared with control group (χ²=13.64, P<0.01), and the severe OSA group had markedly more EPVS than the other...
### Table 2. Analysis of PSG Results Among All Groups

| PSG Index | Mild Group (n=25) | Moderate Group (n=20) | Severe Group (n=18) | Total (n=46) | Control Group (n=44) |
|-----------|------------------|-----------------------|--------------------|-------------|---------------------|
| **AHI** (event/h) | 1.10±0.20 | 2.40±0.20 | 3.50±0.20 | 2.29±0.20 | 1.00±0.20 |
| **LSaO2 (min)** | 8.50±0.20 | 8.10±0.20 | 7.90±0.20 | 8.20±0.20 | 8.50±0.20 |
| **ODI** (event/h) | 1.20±0.20 | 2.30±0.20 | 3.40±0.20 | 2.29±0.20 | 1.00±0.20 |
| **SI** (min) | 2.50±0.20 | 3.60±0.20 | 4.70±0.20 | 3.40±0.20 | 2.50±0.20 |
| **SE (%)** | 85.30±5.20 | 82.30±5.20 | 79.30±5.20 | 81.30±5.20 | 85.30±5.20 |
| **WASO (min)** | 1.50±0.20 | 2.60±0.20 | 3.70±0.20 | 2.40±0.20 | 1.50±0.20 |
| **AI (event/h)** | 1.50±0.20 | 2.60±0.20 | 3.70±0.20 | 2.40±0.20 | 1.50±0.20 |
| **N1%** | 6.50±0.10 | 7.60±0.10 | 8.70±0.10 | 7.60±0.10 | 6.50±0.10 |
| **N2%** | 16.30±0.30 | 17.40±0.30 | 18.50±0.30 | 17.40±0.30 | 16.30±0.30 |
| **N3%** | 11.70±0.50 | 12.80±0.50 | 13.90±0.50 | 12.80±0.50 | 11.70±0.50 |

Notes: P was the p-value produced when comparing OSA group with control group, P1 was the p-value obtained when comparing severe group with mild group, P2 was the p-value obtained when comparing severe group with moderate group. Differences of <0.05 indicated statistical significance.

Abbreviations: AHI, apnea hypopnea index; SE, sleep efficiency; WASO, wake after sleep onset; SI, sleep latency; LSaO2, the lowest blood oxygen saturation; ODI, oxygen desaturation index; N1%, percentage of non-rapid eye movement sleep N1 stage; N2%, percentage of non-rapid eye movement sleep N2 stage; N3%, percentage of non-rapid eye movement sleep N3 stage; REM%, percentage of rapid eye movement sleep. AI, arousal index.

In the present article, our findings suggested that the respiratory parameters AHI, ODI and TS90% were positively correlated with EPVS, while a significant negative correlation was observed between EPVS and LSaO2. Such result suggests that, with the increase in OSA severity, the hypoxia degree in the body during sleep is aggravating, which together with the sleep fragmentation, supports the use of PVS as a vascular clearance impairment marker in sleep disorder. The PVS seen on MRI may not be pathological, but it has been suggested to be related to age-related diseases and the vascular risk factors. On this basis, we speculate that a higher risk for EPVS possibly takes place via the pathophysiologic mechanisms non-specific to the cerebral blood flow, as shown below: (1) the sympathetic activation induced by intermittent hypoxia aggravates systemic inflammatory response, decreases hypocretin secreted by the brain at the time of sleep fragmentation, leads to blood leukocytosis and induces inflammatory response and oxidative damage. The activation of inflammatory response may cause microvascular endothelial cell injury, microvascular wall structural changes and astrocyte swelling, which then leads to interstitial fluid leakage, restricted drainage space and the subsequent PVS development. (2) It is related to the respiratory event-related hemodynamic changes. When the automatic adjustment mechanism of the cerebrovascular system is insufficient to protect the brain against the apnea-related rapid intracranial pressure fluctuations, the hemodynamic changes will result in higher resistance of PVS to CSF clearance, which hinders the elimination of metabolites through the lymphatic system.

Associations between sleep architecture and PVS changes on MRI have also been found in this study, which suggested that SE was significantly negatively correlated with EPVS, and AI was positively correlated with EPVS. The decreased SE may be related to the ineffective PVS drainage, and the lower percentage of time spent asleep is probably related to the ineffective perivascular drainage, which may result in PVS enlargement. Additionally, the noradrenaline produced by the locus coeruleus in the brain is of great significance to promote arousal, which is a key neurotransmitter that regulates the glymphatic system function. The elevation of norepinephrine level may lead to increased cell volume, which
can further reduce the PVS volume fraction and indirectly increase the CSF flow resistance, finally affecting the clearance efficiency of the glymphatic system. On the other hand, the clearance of intracerebral metabolites depends on the polar distribution of astrocyte terminal aquaporin 4 (AQP4). The long-term sleep deprivation or fragmentation affects the distribution of AQP4, renders its loss of polarity and even arrangement on the astrocyte surface, which thereby affects the metabolite elimination efficiency by the glymphatic system, resulting in PVS expansion.

Some important physiological alterations may take place during sleep, as defined by the various sleep stages. However, the sleep-stage distributions may not be measured by SE and AI. Therefore, we further examined the relationships of different sleep-stage proportions with EPVS. As revealed by our findings, EPVS was negatively correlated with N3% and REM%, but positively correlated with N2%. Fultz et al pointed out that the

### Table 3 Partial Spearman’s Rank Order Correlation Results Unadjusted for Covariates

|            | r   | P    |
|------------|-----|------|
| Age        | 0.257 | 0.007 |
| BMI        | 0.298 | 0.002 |
| MMSE       | -0.215 | 0.026 |
| ESS        | 0.296 | 0.002 |
| AHI        | 0.628 | <0.001 |
| LSaO2      | -0.654 | <0.001 |
| TS90%      | 0.585 | <0.001 |
| ODI        | 0.655 | <0.001 |
| SE         | -0.343 | <0.001 |
| AI         | 0.365 | <0.001 |
| N1%        | 0.221 | 0.022 |
| N2%        | 0.621 | <0.001 |
| N3%        | -0.524 | <0.001 |
| REM%       | -0.552 | <0.001 |

**Note:** Differences of P<0.05 indicated statistical significance.

**Abbreviations:** PVS, perivascular space; BMI, body mass index; MMSE, Mini Mental State Scale; ESS, Epworth Sleepiness Scale; AHI, apnea hypopnea index; SE, sleep efficiency; LSaO2, the lowest blood oxygen saturation; TS90% percentage of sleep time spent with an oxygen saturation < 90%; ODI, oxygen desaturation index; N1%, percentage of non-rapid eye movement sleep N1 stage; N2%, percentage of non-rapid eye movement sleep N2 stage; N3%, percentage of non-rapid eye movement sleep N3 stage; REM%, percentage of rapid eye movement sleep; AI, arousal index.

### Table 4 Partial Spearman’s Rank Order Correlation Results Controlling for Age, BMI, ESS

|            | r   | P    |
|------------|-----|------|
| MMSE       | -0.141 | 0.152 |
| AHI        | 0.565 | <0.001 |
| LSaO2      | -0.635 | 0.001 |
| TS90%      | 0.525 | <0.001 |
| ODI        | 0.618 | <0.001 |
| SE         | -0.319 | 0.001 |
| AI         | 0.273 | 0.005 |
| N1%        | 0.101 | 0.310 |
| N2%        | 0.532 | <0.001 |
| N3%        | -0.403 | <0.001 |
| REM%       | -0.478 | <0.001 |

**Note:** Differences of P<0.05 indicated statistical significance.

**Abbreviations:** PVS, perivascular space; BMI, body mass index; MMSE, Mini Mental State Scale; ESS, Epworth Sleepiness Scale; AHI, apnea hypopnea index; SE, sleep efficiency; LSaO2, the lowest blood oxygen saturation; TS90%, percentage of sleep time spent with an oxygen saturation < 90%; ODI, oxygen desaturation index; N1%, percentage of non-rapid eye movement sleep N1 stage; N2%, percentage of non-rapid eye movement sleep N2 stage; N3%, percentage of non-rapid eye movement sleep N3 stage; REM%, percentage of rapid eye movement sleep; AI, arousal index.
lymphatic system was mainly active during sleep to eliminate the tissue waste from the perivascular system and to realize sleep homeostasis, which mainly took place during the phase of slow wave sleep and was triggered or led by the slow brain waves with low activity. Laura Lewis also suggested that CSF flow velocity exceeded the imaging critical velocity (11.4 mm/s for slice 2), which identified a large-amplitude pulsatile flow of CSF at 0.05 Hz that appeared during NREM sleep. Although the glymphatic system was somewhat active during the day, Nedergaard discovered that it was during deep NREM sleep that this sanitation system kicked into high gear. With the pulsing rhythm of deep NREM sleep, the brain expels 10 or 20 times more effluents. At the same time, the volume of brain glial cells shrunk by 60% in deep sleep stage, which created a greater space for the CSF to eliminate the metabolic wastes. However, our results suggested that N2% was positively correlated with EPVS, which might further support the speculation that ISF elimination mainly took place in slow wave sleep process, conforming to research by Xie et al. Moreover, our data suggested that the reduced REM% was associated with EPVS. The REM sleep deprivation is accompanied by the reduced CSF lymphatic circulation, thus resulting in hydrocephalus. Consequently, it is speculated that the EPVS may be related to the mechanical association of the pathological brain status produced by the REM sleep deprivation. In our data, 18 (40.91%) of the control subjects presented EPVS (Table 1), and it was inferred from the data that MRI-visible PVS was considered as the enlarged PVS in our study. Besides, PVS on MRI of normal controls showed normal results, and even their dilation was related to aging.

Notably, in our cohort, MMSE scale was used for evaluating the changes in cognitive function among OSA cases. The results showed that most of the patients with OSA did not have cognitive decline, the score of MMSE was lower in OSA group (27.58±1.39) compared with the control group (28.00±1.29), and the MMSE score of the severe OSA group was significantly lower (27.12±1.58). However, there was no significant difference in the MMSE score between groups, or correlation with EPVS, in contrary to several other reports suggesting that OSA caused different degrees of cognitive dysfunction. This difference might be due to the fact that the MMSE scale was less sensitive than Montreal Cognitive Assessment scale and participants were all young and middle-aged in our study.

Certain limitations should be noted in this study. For instance, this study retrospectively analyzed a small portion of patients diagnosed with OSA at our hospital, which might lead to sampling bias and have limited general applicability. Secondly, PSG serves as the gold standard for sleep staging, but it is only a snapshot of the whole night sleep of an individual subject, which may not represent the long-term sleep mode of a family. Therefore, long-term follow-up studies are warranted to further investigate and explore the role of OSA in the EPVS pathophysiology and whether OSA treatment prevents PVS progression.

As far as we know, no epidemiological studies have reported that OSA is related to EPVS among the young and middle-aged populations. Our study had the main strength that our data represent the general population with a low risk of dementia and cerebrovascular diseases. This study revealed the important insights that the EPVS severity worsened with the increase in sleep disturbances in OSA, suggesting that the severity of OSA might be regulated by the number of pathophysiological mechanisms and affected the outcome of brain structural modification, thus leading to cerebrovascular diseases. Therefore, the increased risk of EPVS in OSA is a potential contributor to the increased risk of stroke in OSA; however, further studies are needed.

Data Sharing Statement
The data that support the findings of this study are available from the corresponding authors upon reasonable request.

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