Sleep related breathing disorders (SRBDs) are common in patients with stroke. However, the relationship between SRBD and stroke is not clear. Central sleep apnea (CSA) is characterized by a loss of central respiratory drive and effort during sleep, resulting in insufficient or absent ventilation, rather than physical upper-airway collapse. CSA is attributed to medical or neurological conditions including stroke. Although brainstem infarctions have previously been suggested to be associated with CSA, the association of lesion location and CSA in patients with ischemic stroke has not been well elucidated. A 69-year-old man with a history of hypertension and diabetes mellitus was admitted due to stroke. The brain magnetic resonance imaging showed an acute ischemic stroke in the right ventral thalamus and adjacent hypothalamus. During hospitalization, polysomnography (PSG) was performed because repetitive cessation of respiration during sleep was observed by chance. PSG showed severe CSA; the apnea-hypopnea index (AHI) was 73.5 with a minimum oxygen saturation of 89% and central apnea index (CAI) was 63.0. Two years later, follow-up PSG showed that AHI was 7.2 with a minimum oxygen saturation of 91% and CAI was 1.0.

We report the patient with CSA after ischemic stroke with right thalamus and adjacent hypothalamus, which resolved spontaneously with time.

### Key Words:
Stroke, Central sleep apnea, Polysomnography.

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Central Sleep Apnea after Stroke

stroke and follow up PSG showed that AH1 was 7.2 with a minimum oxygen saturation of 91% and CAI was 1.0 (Table 1).

Discussion

Although obstructive apnea is more common, central apnea may be noted initially after stroke. It is difficult to determine the relationship between prior and post-stroke SRBD in case of no prior objective diagnosis of SRBD. Although our case did not have PSG finding before stroke onset, it is reasonable to assume that CSA caused by acute ischemic stroke because it resolved spontaneously over two years. The CSA following a stroke is abrupt in onset and tend to resolve with time. However, well-illustrated PSG finding showing typical CSA without CSB has rarely been reported. On this case, the CAI was normalized and markedly reduced compared to initial PSG (63.0 vs. 1.0).

This case showed acute ischemic lesion in the right ventral thalamus and adjacent hypothalamus. Post-stroke sleep-disordered breathing has been well introduced but no specific lesion location was related to the SRBD as well as CSA. Previous studies showed that brainstem stroke caused CSB and

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**Figure 1.** Diffusion weighted imaging of the patient. Axial view (A) and coronal view (B) shows acute ischemic stroke in the right ventral thalamus and adjacent hypothalamus.

**Figure 2.** Polysomnographic (PSG) finding 3 days after stroke. A 5-minutes epoch PSG shows periodic central respiratory events without Cheyne-Stokes breathing. The overall apnea-hypopnea index was 73.5 and central apnea index was 63.0. Two electrooculogram channels (LOC-M2, ROC-M1), two chin electromyogram channels (Chin-Chin1, Chin1-Chin2), and six electroencephalogram channels (F3-M2, F4-M1, C3-M2, C4-M1, O1-M2, O2-M1) were measured. LAT: left anterior tibialis electromyogram, RAT: right anterior tibialis electromyogram, Snore: snore sensor, CHEST: chest respiratory movement, ABD: abdomen respiratory movement, PTAF: pressure transducer air-flow sensor, Thermister: oro-nasal thermal sensor, EKG: electrocardiogram, SaO2: oxyhemoglobin saturation by pulse oximetry, HR: heart rate, CA: central apnea.
high prevalence of CSA without CSB. In a few additional reports, various lesions such as cortical and subcortical area have been reported to be associated with central periodic breathing during sleep in acute ischemic stroke. The mechanism of CSA after stroke has been discussed to be a direct consequence of the injury of central nervous system structures, which is involving autonomic and volitional respiratory centers. Within the brainstem, the medulla oblongata contains central chemoreceptors that response to a loop-gain feedback system on the PaCO₂. Lesions to this area lead to decrease chemosensitivity during sleep and develop CSA. Besides, non-brainstem regions might have role of autonomic and voluntary respiration involving insular, and supplementary motor cortex and thalamus. It is also known that some hypothalamic nuclei are interconnected with respiratory nuclei located in the brainstem and involve respiratory control as well as other autonomic brain function. Although dysfunction of hypothalamus causes abnormal breathing, the role of the hypothalamus in regulation of respiration during sleep remain to be elucidated. Similarly, it is not well known yet about specific anatomical structures and pathomechanism related to CSA after stroke. This case demonstrates an example of CSA without CSB after ischemic stroke with right thalamus and adjacent hypothalamus, which resolved spontaneously with time. Further elucidation of mechanisms how these structures regulates involuntary respiratory function during sleep is required.

Conflicts of Interest

The authors have no potential conflicts of interest to disclose.

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| Table 1. Polysomnographic parameters between initial and follow-up study |
|--------------------------|--------------------------|
|                          | Initial PSG              | PSG after 2 years |
| Total recording time, min | 496.5                    | 441              |
| Total sleep time, min    | 264.5                    | 301.5            |
| Total stage N1, min      | 121 (24.4%)              | 55.5 (12.6%)     |
| Total stage N2, min      | 143.5 (28.9%)            | 195.5 (44.3%)    |
| Total stage N3, min      | 0                        | 0                |
| Total stage R, min       | 0                        | 50.5 (11.5%)     |
| Wake after sleep onset, min | 230.5                   | 119.5            |
| Latency to sleep onset, min | 1.5                      | 20               |
| Latency to REM sleep, min | NC                       | 102              |
| Sleep efficiency, min    | 53.3%                    | 68.4%            |
| Apnea-hypopnea index     | 68.7                     | 7.2              |
| Central apnea index      | 63.0                     | 1.0              |
| Nadir oxygen saturation  | 89%                      | 91%              |

PSG: polysomnography, REM: rapid eye movement, NC: no contribution