Sleepiness after Stroke: Case Report and Review of Literature on Hypersomnia as a Result of Stroke

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Introduction

Daytime sleepiness is defined as the inability to stay awake and alert during the major waking episode of the day resulting in irrepressible need for sleep or unintended lapses into drowsiness. Excessive sleepiness affects up to 5% of the general population and can result in significant disability. Hypersomnia can be either the result of unmet sleep needs or can also be due to reduced activity of alerting systems in the brain. The former can be due to sleep restriction or from sleep disruption. The International Classification of Sleep disorders categorize various disorders of hyper somnolence into different sub-groups. Exaggerated sleep propensity with daytime sleepiness and/or prolonged nocturnal sleep following cerebrovascular accident falls under ‘Hypersomnia due to a medical disorder’. Sleepiness after stroke is a rare complication but when it happens, can be disabling to the patients. Hypersomnia may have worse outcomes than apathy [1] in stroke patients.

The exact frequency of hypersomnia after stroke is unknown though in a few studies, it is seen up to 1.1 to 27% of the patients. The severity of sleepiness following stroke can be very mild or can be severe and can be transient or permanent. The location of the stroke and in some cases, the size of the stroke is the major determinant of the nature of the hypersomnia that follows the stroke. In this article, we describe a case of an elderly female who developed severe sleepiness after having a stroke in bilateral Para median thalamic region and also review the literature on sleepiness as a delayed complication of stroke.

Case Report

An 87-year-old female was taken to an outside hospital for acute onset of somnolence. Her past medical history is significant for hypertension and diabetes mellitus. Magnetic resonance imaging (MRI) of the brain performed at the outside facility showed the presence of bilateral thalamic and sub-thalamic infarctions (Figure 1). Magnetic resonance angiography (MRA) did not reveal anomalous vasculature in the posterior circulation though the patient did not undergo conventional angiogram at that time. The patient was drowsy for the first few days but improved gradually during the hospital stay. During the hospital course, both the patients and the relatives reported...
significant reduction in her hand tremor for which she had been on propranolol. She started to experience sleepiness in the next few weeks, which persisted subsequently.

Historical Aspects


Stroke and its Relationship with Sleep and Sleep Disorders

Sleep has protective effect against strokes and hence the incidence of stroke increases in sleep disorders. Sleep wake cycle is regulated by a complex interaction of various different areas of the brain like brainstem, hypothalamus, thalamus and preoptic region. This is why focal lesions in these regions result in sleep disorders. Stroke patients who experience sleep disorders can range anywhere from 20 to 63% [6]. A wide range of sleep disorders can be seen in stroke patients including hypoponsomnia, insomnia, parasomnias, periodic limb movements in sleep (PLMS) and sleep related breathing disorders. The presence of sleep disorders in stroke patients may be associated with increased mortality and poorer prognosis [7]. Strokes can also affect the sleep architecture. A stroke in the supratentorial region has been linked with decrease in total sleep time with a reduction of Non Rapid Eye Movement (NREM) sleep [8]. There may be ipsilateral or bilateral reduction in sleep spindles after a hemispheric stroke [9]. Temporary reduction in Rapid Eye Movement (REM) sleep may also be observed in some cases of supratentorial stroke. This is more frequent with right-sided strokes. The saw tooth wave may be reduced after hemispheric stroke [10]. Occipital stroke can reduce REM sleep [11]. Strokes in pontomesencephalic junction and the Raphe nucleus may diminish NREM sleep while not affecting the amount of REM sleep. There may be REM sleep onset at the beginning of the sleep cycle. At the same time, strokes in lower pons may reduce REM sleep selectively [12-15]. Para median thalamus and lower pontine strokes can result in lack of slow wave sleep and preserved REM sleep. Ponto medullary junction and mesencephalic tegmental strokes may increase REM sleep [6] whereas infarcts in the midbrain can result in increases in both NREM and REM sleep [16].

Insomnia is common in stroke patients and affects 20-56% of stroke patients (higher incidence than hypoponsomnia). About 18% of people reported new onset insomnia after the stroke. Insomnia typically occurs in the acute phases of the stroke. Thalamic and brainstem (thalamomesencephalic, pontomesencephalic and pontine tegmental) strokes can present with insomnia [7,17].

Various parasomnias like REM sleep behavior sleep disorders (RBD) have been described with tegmental pontine strokes, strokes in the thalamus and in the temporal, parietal, and occipital lobes. In one case report, pontine stroke was associated with isolated cataplexy and REM sleep behavior disorder [18].

Denovo leg syndromes were reported to be as high as 12% in all stroke patients [19]. Restless leg syndrome was reported in strokes involving pons, thalamus, basal ganglia and corona radiata. About 2/3rd of those restless leg syndrome patients reported bilateral symptoms while 1/3rd reported symptoms on the side opposite to the stroke [6].

Sleep disordered breathing (SDB) can be seen in as high as 50 to 70% in stroke patients [20]. Not only does SDB increase the risk for stroke, stroke also can result SDB. Both obstructive sleep apnea and central sleep apnea have been reported widely after stroke. Identifying the occurrence of sleep-disordered breathing is an important component of primary or secondary stroke prevention.

The most severe cases of hyper somnolence secondary to stroke are caused by Para median thalamic infarctions. A case of Kleine-Levin syndrome was also reported after multiple cerebral infarctions [21]. Narcolepsy with low CSF hypocretin levels was reported in a case after diencephalic stroke [22]. Circadian rhythm disturbances were noted after modafinil treatment in a patient with bilateral thalamic stroke related sleepiness [23].

Mechanism of Hypoponsomnia after Stroke

A reduction in the activity of the arousal system (dearousal) underlies most forms of post stroke hypoponsomnia [24]. Bilateral lesions of posterior hypothalamic, tegmental midbrain and
upper pons tend to cause severe forms of hyper somnolence as fibers of arousal systems are bundled and can be affected by a single lesion. The mental arousal seems more affected by medial lesions whereas the lateral lesions tend to affect motor arousal preferentially [25]. Other areas that may occasionally cause hypersomnia are stratum, pontine tegmentum, medulla and cerebral hemisphere (if large area is involved). It is seen that hemisphere strokes resulting in hypersomnia tend to be more on the left side than right and anterior more than posterior. Cortical and subcortical strokes can also cause post stroke hypersomnia but with wide variability which may be related to a relatively less compact arousal pathways with respect to the brainstem.

Conclusion
Sleepiness following stroke is an under-recognized complication and can affect up to a quarter of stroke patients with a severity ranging from mild to very severe. Patients with hypersomnia after stroke are 10 times more likely go to nursing home compared to those who do not have hypersomnia. The recovery from hypersomnia is very variable with most cases improving in the first few days. However, hypersomnia can persist for years and can evolve into akinetic mutism. Therapy is often ineffective [6] in the patients who are affected severely. Multiple medications including amphetamine salts, modafinil, methylphenidate, levodopa, bromocriptine have been tried with variable success. Prompt recognition of this complication and a thorough sleep evaluation can lower the morbidity and improve patient outcomes [26].

References